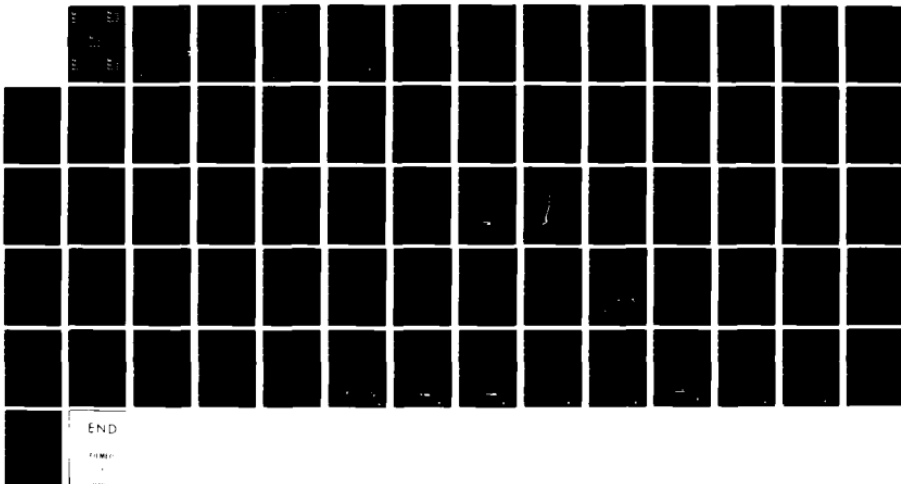


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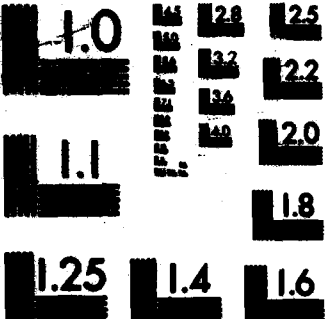
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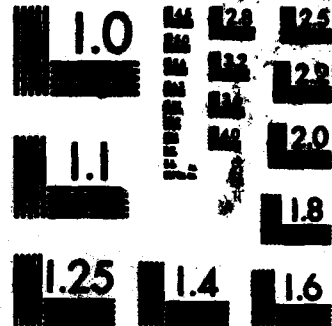


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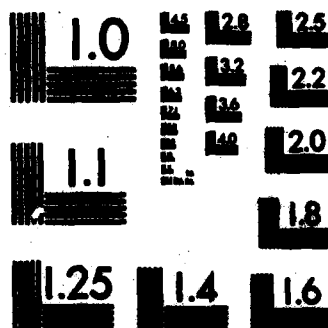
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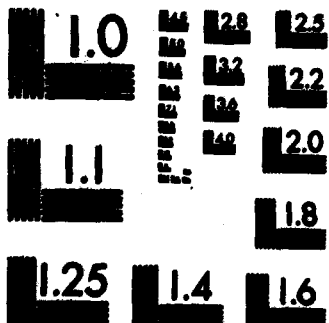
MICROCOPY RESOLUTION TEST CHART
NATIONAL BUREAU OF STANDARDS-1963-A



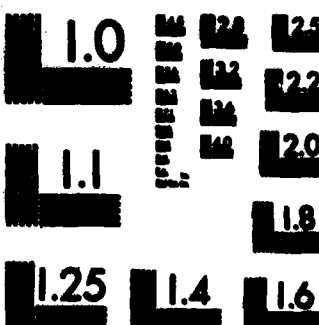
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MICROCOPY RESOLUTION TEST CHART
NATIONAL BUREAU OF STANDARDS-1963-A

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REPORT DRXTH-TE-CR-2135

(12)

**CHEMICAL ANALYSIS SUPPORT: LIMITED
ANALYSIS OF BRISTOL, RHODE ISLAND WELL
WATER SAMPLES**

Kevin Beltis
Christine Jones
Linda Sadowski

ARTHUR D. LITTLE, INC.
CAMBRIDGE, MA 02140

SEPTEMBER, 1982

FINAL TASK REPORT

DISTRIBUTION UNLIMITED: APPROVED
FOR PUBLIC RELEASE



prepared for

U.S. Army Toxic and Hazardous Material Agency,
Aberdeen Proving Ground, Maryland 21010

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SECURITY CLASSIFICATION OF THIS PAGE (When Data Entered)

REPORT DOCUMENTATION PAGE		READ INSTRUCTIONS BEFORE COMPLETING FORM
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		6. PERFORMING ORG. REPORT NUMBER
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18. SUPPLEMENTARY NOTES		
19. (If necessary, enter data on separate sheet and identify by block number) Volatile, Acid, Base/Neutral and Pesticide/PCB Priority Pollutants 1,1-dimethyl-hydrazine Bristol, Rhode Island		
20. (If necessary, enter data on separate sheet and identify by block number) Well water samples collected from Bristol, Rhode Island were analyzed for volatile, acid, base/neutral and pesticide/PCB priority pollutants by the Federal Register Methods. Detection limits for the methods were determined following the May, 1980 USATHAMA QA program. In addition, attempts were made to detect the presence of UDMH in these field samples.		

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Appendix C - Priority Pollutant Base-Neutral Data	
Appendix D - Priority Pollutant Pesticide/PCB Data	

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SUMMARY

Field samples collected for this task were analyzed for volatile, acid, base-neutral and pesticide priority pollutants using EPA methods. None of these pollutants were detected above the method detection limits determined from quality control samples.

In addition, experiments to develop an analytical method for 1,1-dimethyl-hydrazine were conducted but no satisfactory method was found in the time allotted. However, based on extraction and detection characteristics of similar compounds, it is estimated that 1 ppm of UDMH, if present would have been detected by gas chromatography/mass spectrometry analysis.

I. Sampling

On September 9, 1980 an employee of Arthur D. Little, Inc., met with USATHAMA personnel in Bristol, Rhode Island to assist in the collection of well water samples. Two distinct samples were collected, each consisting of two, 40-mL serum vials for the volatile organic analysis and two, gallon glass jars with teflon lined screw caps for the remaining analyses. Of the two samples, one represented the lower, aqueous phase present in the well, while the second sample represented the upper, oily phase present in the well. Table I-1 lists the samples collected and their subsequent laboratory sample numbers.

Table I-1

Samples Collected at Bristol Site

<u>Field Sample Number</u>	<u>Description</u>	<u>Laboratory Number</u>
BR 100	2 gallons, aqueous phase	BR 100 (composite)
BR 102		
BR 101	45-mL volatile organics, duplicate samples	BR 101
BR 103	aqueous phase	(BR 103) ¹
BR 104	2 gallons, ~10% oil	BR 104 (composite)
BR 105		
BR 106	45-mL volatile organics, duplicate samples	(BR 106) ¹
BR 107	oily phase	BR 107

¹Duplicate samples not analyzed

II. Sample Preparation/Analysis

Samples for the volatile organic analysis require no sample preparation and are analyzed directly by the purge and trap method¹ (Federal Register, Purgeables Method 624). One liter aliquots of BR 100 were extracted for acid and base-neutral analyses (Method 625) after initial pH adjustment, with three 60-mL volumes of methylene chloride. After drying each extract over sodium sulfate, it was concentrated to a final volume of 2-mL by Kuderna-Danish. Ten microliters of a d₁₀-anthracene standard solution was added as an internal standard just prior to analysis by gas chromatography/mass spectrometry¹ (Federal Register, Base-Neutrals Method 625).

A one-liter aliquot of BR 100 was also extracted for the priority pollutant pesticide analysis with three 60-mL volumes of 15 percent methylene chloride in hexane. The extract was concentrated to less than 10 mL and eluted through Florisil using 6 percent, 15 percent, and 50 percent diethylether in petroleum ether. After final concentration of each fraction to 10 mL, they were analyzed by gas chromatography using an electron capture detector². In addition, 1 mL of BR 104, the oil phase, was diluted to 10 mL with hexane and analyzed by GC/ECD. No characteristic pesticide/PCB patterns were observed with this initial screen, making Florisil fractionation unnecessary.

Several attempts were made to analyze samples for unsymmetrical dimethylhydrazine (UDMH). An analytical standard (99%+ pure) was obtained from Aldrich Chemical Company (Cat. No. D16, 1608). Of the analytical methods cited in the literature, one was examined³. This method involves derivatization (with acetone), solvent extraction (pH adjustment with Base and CH₂Cl₂ extraction) and analysis by gas chromatography using a flame ionization detector. Laboratory samples were prepared by spiking neat UDMH into standard water (100 mg/L sulfate and chloride in distilled water) in the range of 0.1 to 10 parts per million. These spikes, along with aliquots of the two field samples, were analyzed as outlined in the method. No UDMH was detected from any of the samples analyzed. In addition to the OV-101 column recommended in the method, a 10% SP-2100 and a 5% SP-1000 column were examined. Column temperatures from 75°C to 200°C were tried in an effort to resolve the derivative, if present, from the solvent front. None of the conditions tried proved successful.

1,2-Diphenylhydrazine, one of the base-neutral priority pollutants, is routinely analyzed in our laboratories by GC/MS, with a calculated detection limit of 15 ppb (see Appendix C). Hydrazine can be reproducibly detected in the range of 30-50 ppb. Based on these data, it is estimated that < 1 ppm UDMH would have been detected by GC/MS if it were present in the extracted sample.

In addition, the Bristol samples were examined based on mass spectral data. No evidence of UDMH was found in any of the three fractions examined (acid, base/neutral and volatiles).

III. Quality Control

The ability of Arthur D. Little, Inc. to perform priority pollutant analyses on samples for USATHAMA has been well documented in Tasks R902.35.03, R902.35.08, and R902.35.12. During the period of time the Bristol samples were being analyzed, Quality Control data for the priority pollutants were being generated under Task R902.35.08 of this program. Because of the Quick Response nature of the program and the limited number of samples that were to be analyzed for the Bristol site, authorization was given to reference this data for Quality Control purposes during sample analysis. Field samples obtained for Task R902.35.08 were spiked at five concentration levels and analyzed by the Federal Register methods. The detection limits were determined as stipulated in the May 1980 USATHAMA QA Program. These data are shown in the Appendices, including the slope (n), the correlation coefficient (R), and the detection limit at the 90% confidence level [$x(d)$], calculated by the method of Hubaux and Vos.

In addition, during analyses of the Bristol samples, samples were being analyzed concurrently for an EPA-sponsored contract. The quality control data from these EPA analyses are included in this report to demonstrate the fact that the laboratory and method were operating in control during analyses of the Bristol samples.

IV. RESULTS

The results of the analysis of the Bristol samples for volatile priority pollutants can be found in Appendix A. The total ion and reconstructed ion chromatograms of the Bristol sample show that only the internal standards are detected. These were added prior to sample analysis. The corresponding chromatograms for acid and base/neutral priority pollutant analyses reflect the character of the water sampled. The well from which the samples were obtained contained substantial quantities of an oil type material as described in the sampling section. The presence of this material provided a background spectra illustrated in the sample chromatograms for the acid and base/neutral fractions. The detection limits reported for the acid and base/neutral priority pollutants were generated under a separate ongoing task of this program as described in Section III. Because of the possibility of interferences from the hydrocarbons in the Bristol samples, the detection limits may be higher than the x(d) values reported for the quality control samples.

The results of the analysis of the florisil fractions of the Bristol samples indicate that no pesticide or PCB priority pollutants were detected at a level higher than the calculated detection limits.

V. References and Footnotes

¹Federal Register 12/3/79. Vol. 44, No. 233. Purgeables Method 624, Base/Neutrals, Acids Method 625.

²U.S. EPA. Evaluation of Protocols for Pesticides and PCB's in Raw Wastewater. EPA-600/2-79-166.

³J. T. Veal. "The Analysis of Hydrazine, Monomethyl Hydrazine and 1,1-Dimethyl Hydrazine Using WCOT/GC Techniques." Air Force Syst. Command, Civ. Environ. Eng. Dev. Off., (Tech. Rep.) CEEDO-TR (U.S.), No. CEEDO-TR-78-14, Proc. Conf. Environ. Chem. Hydrazine Fuels, 1977, p 79-98 (1978).

APPENDIX A

Method for the Analysis of Organic Volatiles in Water

Priority Pollutant Volatile Organics Data

ANALYSIS OF ORGANIC VOLATILES
IN WATER

APPLICATION

METHOD USED TO DETERMINE THE CONCENTRATIONS OF THE FOLLOWING COMPOUNDS IN WATER SAMPLES;

CHLOROMETHANE
DICHLORODIFLUOROMETHANE
BROMOMETHANE
VINYL CHLORIDE
CHLOROETHANE
METHYLENE CHLORIDE
TRICHLOROFLUOROMETHANE
1,1-DICHLOROETHYLENE
1,1-DICHLOROETHANE
TRANS-1,2-DICHLOROETHYLENE
CHLOROFORM
1,2-DICHLOROETHANE
1,1,1-TRICHLOROETHANE
CARBON TETRACHLORIDE
BROMODICHLOROMETHANE
1,2-DICHLOROPROPANE
TRANS-1,3-DICHLOROPROPYLENE
TRICHLOROETHYLENE
BENZENE
CIS-1,3-DICHLOROPROPYLENE
DIBROMOCHLOROMETHANE
1,1,2-TRICHLOROETHANE
Bromoform
1,1,2,2-TETRACHLOROETHANE

1,1,2,2-TETRACHLOROETHYLENE

TOLUENE

CHLOROBENZENE

ETHYL BENZENE

A. TESTED CONCENTRATION RANGE:
5-50 UGL FOR EACH

B. SENSITIVITY; SEE FEDERAL REGISTER VOL. 44, NO. 233.

C. DETECTION LIMIT:

CHLOROMETHANE	18 UGL
DICHLORODIFLUOROMETHANE	42 UGL
BROMOMETHANE	15 UGL
VINYL CHLORIDE	21 UGL
CHLOROETHANE	28 UGL
METHYLENE CHLORIDE	18 UGL
TRICHLOROFLUOROMETHANE	14 UGL
1,1-DICHLOROETHYLENE	18 UGL
1,1-DICHLOROETHANE	13 UGL
TRANS-1,2-DICHLOROETHYLENE	14 UGL
CHLOROFORM	10 UGL
1,2-DICHLOROETHANE	10 UGL
1,1,1-TRICHLOROETHANE	17 UGL
CARBON TETRACHLORIDE	15 UGL
BROMODICHLOROMETHANE	11 UGL
1,2-DICHLOROPROPANE	11 UGL
TRANS-1,3-DICHLOROPROPYLENE	10 UGL
TRICHLOROETHYLENE	13 UGL
BENZENE	12 UGL
CIS-1,3-DICHLOROPROPYLENE	8 UGL
DIBROMOCHLOROMETHANE	9 UGL
1,1,2-TRICHLOROETHANE	11 UGL
BROMOFORM	9 UGL
1,1,2,2-TETRACHLOROETHANE	9 UGL
1,1,2,2-TETRACHLOROETHYLENE	11 UGL

TOLUENE	12 UGL
CHLOROBENZENE	12 UGL
ETHYL BENZENE	12 UGL

D. INTERFERENCES; SEE FEDERAL REGISTER 12/3/79
VOL. 44, NO. 233

E. ANALYSIS RATE; AFTER TUNING OF INSTRUMENT, ONE
ANALYST CAN ANALYZE 6-7 FIELD SAMPLES IN AN
8-HOUR DAY

CHEMISTRY

CH₃Cl CHLOROMETHANE
CAS RN 74-87-3
MELTING PT - 97C BOILING PT - 23.7C

CCL₂F₂ DICHLORODIFLUOROMETHANE
CAS RN 75-71-8
MELTING PT - 158C BOILING PT - 29.8C

CH₃Br BROMOMETHANE
CAS RN 74-83-9
MELTING PT - 93.66C BOILING PT 3.56C

C₂H₃Cl VINYL CHLORIDE
CAS RN 9003-22-9
MELTING PT - 160C BOILING PT - 14C

C₂H₅Cl CHLOROETHANE
CAS RN 75-00-3
MELTING PT - 138.7C BOILING PT 12.3C

CH₂Cl₂ METHYLENE CHLORIDE
CAS RN 75-09-2
MELTING PT - 97C BOILING POINT 39.8-40C

CCL₃F TRICHLOROFLUOROMETHANE
CAS RN 75-69-4
MELTING PT - 111C BOILING PT 23.7C

C₂H₃CL₂ 1,1-DICHLOROETHYLENE
CAS RN 75-35-4
MELTING PT - 122C BOILING PT 30-32C

C₂H₄CL₂ 1,1-DICHLOROETHANE
CAS RN 75-34-3
MELTING PT - 97C BOILING PT 57C

C₂H₂CL₂ TRANS-1,2-DICHLOROETHYLENE
CAS RN 156-60-5
MELTING PT - 50C BOILING PT 48C

CHCL₃ CHLOROFORM
CAS RN 865-49-6
MELTING PT - 63.5C BOILING PT 60.3-61.5C

C₂H₄CL₂ 1,2-DICHLOROETHANE
CAS RN 107-06-2
MELTING PT - 35C BOILING PT 83C

C₂H₃CL₃ 1,1,1-TRICHLOROETHANE
CAS RN 71-55-6
MELTING PT - 35C BOILING PT 74-76C

CCL₄ CARBON TETRACHLORIDE
CAS RN 56-23-5
MELTING PT - 23C BOILING PT 77C

CHBRCL₂ BROMODICHLOROMETHANE
CAS RN 75-27-4
MELTING PT - 55C BOILING PT 87C

C₃H₆CL₂ 1,2-DICHLOROPROPANE
CAS RN 78-87-5
MELTING PT - 100C BOILING PT 95-96C

C₃H₄CL₂ TRANS-1,3-DICHLOROPROPYLENE
CAS RN 78-88-6
BOILING PT 112C

C₂HCL₃ TRICHLOROETHYLENE
CAS RN 79-01-6
MELTING PT - 87C BOILING PT 87C

C₆H₆ BENZENE
CAS RN 71-43-2
MELTING PT 5.5C BOILING PT 80.2C

C3H4CL₂ CIS-1,3-DICHLOROPROPYLENE
CAS RN 78-88-6
BOILING PT 104.3C

CHBR₂CL DIBROMOCHLOROMETHANE
CAS RN 124-48-1
MELTING PT - 22C BOILING PT 119-120C

C₂H₃CL₃ 1,1,2-TRICHLOROETHANE
CAS RN 79-00-5
MELTING PT - 37C BOILING PT 110-115C

CHBR₃ BROMOFORM
CAS RN 75-25-2
MELTING PT 8C BOILING PT 146-150C

C₂H₂CL₄ 1,1,2,2-TETRACHLOROETHANE
CAS RN 79-34-5
MELTING PT - 43C BOILING PT 147C

C₂CL₄ 1,1,2,2-TETRACHLOROETHYLENE
CAS RN 127-18-4
MELTING PT - 22C BOILING PT 121C

C₇H₈ TOLUENE
CAS RN 108-88-3
MELTING PT - 93C BOILING PT 111C

C₆H₅CL CHLOROBENZENE
CAS RN 108-90-7
MELTING PT - 46C BOILING PT 132C

C₈H₁₀ ETHYL BENZENE
CAS RN 100-41-4
MELTING PT - 95C BOILING PT 136C
USE CAUTION IN HANDLING. POTENTIAL TOXIC
INHALATION, AND SKIN ABSORPTION HAZARDS EXIST.

APPARATUS

A. INSTRUMENTATION;
PURGE AND TRAP - TEKMAR MODEL LSC-1
LIQUID SAMPLE CONCENTRATOR

GAS CHROMATOGRAPH - FINNIGAN 4000 GAS
CHROMATOGRAPH/MASS SPECTROMETER WITH A
6110 DATA SYSTEM

B. PARAMETERS:

AMBIENT TEMPERATURE TRAP - 8" OF STAINLESS STEEL TUBING PACKED WITH 0.3" 3% OV-1: 5" TENAX GC; 2.7" SILICA GEL AND GLASS WOOL.

COLUMN - 0.2% CARBOWAX 1500 ON 80/100 CARBOPACK PACKED IN 6-FT X 2 MM ID GLASS COLUMN

GAS FLOW - HELIUM AT 40 ML/MIN

TEMPERATURE -

TRAP - 180C

COLUMN - INITIAL 60C FOR 3 MIN PROGRAM TO 165C AT 8C/MIN. HOLD AT 165C FOR 15 MIN.

INJECTION VOLUME - 5 ML PURGED

DETECTOR - MASS SPECTROMETER: ELECTRON IMPACT DATA ACQUIRED OVER MASS RANGE 21-260 AMU, SCANNING EVERY 2 SECONDS

RETENTION TIME - SEE FEDERAL REGISTER 12/3/79 Vol. 44, NO. 233.

C. HARDWARE/GLASSWARE; SEE FEDERAL REGISTER 12/3/79 VOL. 44, NO. 233.

D. CHEMICALS; SEE FEDERAL REGISTER 12/3/79 VOL. 44, NO. 233.

SUPELCO INC. PURGABLE A, CATALOG #4-8815,
0.2 MG/ML CONCENTRATION

SUPELCO INC. PURGABLE B, CATALOG #4-8816,
0.2 MG/ML CONCENTRATION

SUPELCO INC. PURGABLE C, CATALOG #4-8817,
0.2 MG/ML CONCENTRATION

SUPELCO INC. INTERNAL STANDARD, CATALOG #4-8823, 20 MG/ML CONCENTRATION

STANDARDS

A. CALIBRATION STANDARDS

TALK 1 ML EACH OF SUPELCO, INC. PURGEABLES A, B AND C (SEE SECTION APPARATUS D) AND 1 ML OF METHANOL TO MAKE MASTER SPIKE SOLUTION AT 50 UG/ML CONCENTRATIONS. PREPARE WORKING CALIBRATION STANDARDS USING MILLI-Q WATER AS FOLLOWS.

<u>CAL</u>	<u>ML STOCK/2 ML</u>	<u>FINAL CONC.</u>
CAL 1	2 ML	50 UG/ML
CAL 2	1 ML	25 UG/ML
CAL 3	.5 ML	12.5 UG/ML
CAL 4	.2 ML	5 UG/ML

NEW STANDARDS ARE PREPARED FOR EVERY VOA SET WHICH IS ANALYZED OR AT A MINIMUM EVERY TWO WEEKS. THE STANDARDS ARE STORED IN SEALED VIALS WHICH ARE PACKED IN PAINT CANS FILLED WITH ACTIVATED CHARCOAL AND REFRIGERATED. IN ADDITION, SPIKE ALL CALIBRATION STANDARDS AND FIELD SAMPLES WITH SUPELCO INTERNAL STANDARD STOCK AS FOLLOWS:

INTERNAL STANDARD STOCK = 20 MG/ML
BROMOCHLOROMETHANE
20 MG/ML 1,4-
DICHLOROBUTANE
20 MG/ML 1-CHLORO-
2-BROMO PROPANE

TAKE 10 UL OF INTERNAL STANDARD STOCK AND DILUTE TO 10 MLS WITH METHANOL TO MAKE A MASTER INTERNAL STANDARD SPIKING STOCK AT 20 MG/L.

SPIKE 5 ML SAMPLES WITH 5 UL MASTER INTERNAL STANDARD SPIKING STOCK TO GIVE FINAL CONCENTRATION OF 20 UGL.

SPIKE 2 ML CALIBRATION STANDARDS WITH 2 UL MASTER INTERNAL STANDARD SPIKING STOCK TO GIVE FINAL CONCENTRATION OF 20 UGL.

B. CONTROL SPIKES;
PREPARE THE SAME WAY AS THE CALIBRATION
STANDARDS ONLY PREPARE IN 2 ML NATURAL
WATER

PROCEDURE

SEE FEDERAL REGISTER 12/3/79 VOL. 44, NO. 233.

CALCULATIONS

USING CONTROL SPIKE SAMPLE CONCENTRATION AS CALCULATED FROM DAILY CALIBRATION, PLOT UGL ADDED VERSUS UGL FOUND, BY THE METHOD OF HUBAUX AND VOS, USING THE DETECTION LIMIT TAPE SUPPLIED BY USATHAMA. CORRECT ALL FIELD SAMPLE CONCENTRATIONS USING THE SLOPE OF THE LINEAR REGRESSION LINE.

TABLE A-1

QC NATURAL WATER SPIKES^a
 Task R 902.35.08
 PRIORITY POLLUTANT CHEMICAL ANALYSIS (µg/L)
 PURGEABLE (VOLATILES)

SPIKE LEVEL		+5	+12.5	+20	+25	+50	R	X _d
Chloromethane		—	4.7	NA	19.2	36.9	.991	18.1
Dichlorodifluoromethane		3.8	12.5	NA	23.2	33.7	.974	42.3
Bromomethane		6.9	12.5	NA	22.9	37.5	.996	14.7
Vinyl chloride		7.4	14.4	NA	23.6	36.9	.993	20.6
Chloroethane		6.9	12.0	NA	26.3	39.9	.986	28.4
Methylene chloride		3.7	7.9	10.3	15.3	23.6	.988	17.6
Trichlorofluoromethane		6.4	13.9	19.1	25.7	41.1	.993	13.6
1,1-Dichloroethylene		6.4	13.6	NA	25.3	41.2	.983	18.4
1,1-Dichloroethane		5.7	14.5	18.2	28.6	50.8	.993	12.9
Trans-1,2-dichloroethylene		5.3	13.7	16.8	26.5	46.8	.993	13.5
Chloroform		4.5	12.7	17.1	25.2	45.6	.996	9.9
1,2-Dichloroethene		5.3	14.9	18.8	26.7	49.2	.996	9.7
1,1,1-Trichloroethane		5.6	12.8	14.7	24.8	42.4	.988	17.2
Carbon tetrachloride		5.6	12.2	NA	23.4	39.5	.996	15.0
Bromodichloromethane		5.0	11.0	NA	22.2	39.1	.998	11.4
1,2-Dichloropropene		6.5	14.7	18.6	26.8	46.9	.996	10.6
Trans-1,3-dichloropropene		5.2	13.0	17.2	25.6	48.1	.996	9.5
Trichloroethylene		5.2	12.4	15.2	23.6	42.4	.994	12.6
Benzene		4.4	13.3	17.2	27.3	49.3	.994	12.5
Cis-1,3-dichloropropene		5.0	12.1	NA	24.5	45.3	.999	7.6
Dibromoethane		4.5	10.2	NA	21.3	38.8	.999	9.1
1,1,2-Trichloroethane		4.7	13.3	NA	25.0	45.4	.998	11.4
Bromoform		4.2	8.6	13.1	18.5	38.9	.997	9.1
1,1,2,2-Tetrachloroethene		4.8	13.6	18.1	26.6	49.8	.997	9.1
1,1,2,2-Tetrachloroethylene		5.0	12.0	NA	23.2	41.4	.998	10.6
Toluene		5.1	13.2	17.3	27.2	50.1	.995	11.6
Chlorobenzene		5.5	13.2	16.7	26.3	47.5	.994	12.5
Ethyl benzene		5.3	13.4	17.1	27.2	49.8	.994	12.4

NA = Not Analyzed

^aField sample from uncontaminated background well bordering a U.S. Army installation site.

TABLE A-2
EPA QUALITY CONTROL DATA SUMMARY
PURGEABLE (VOLATILES)

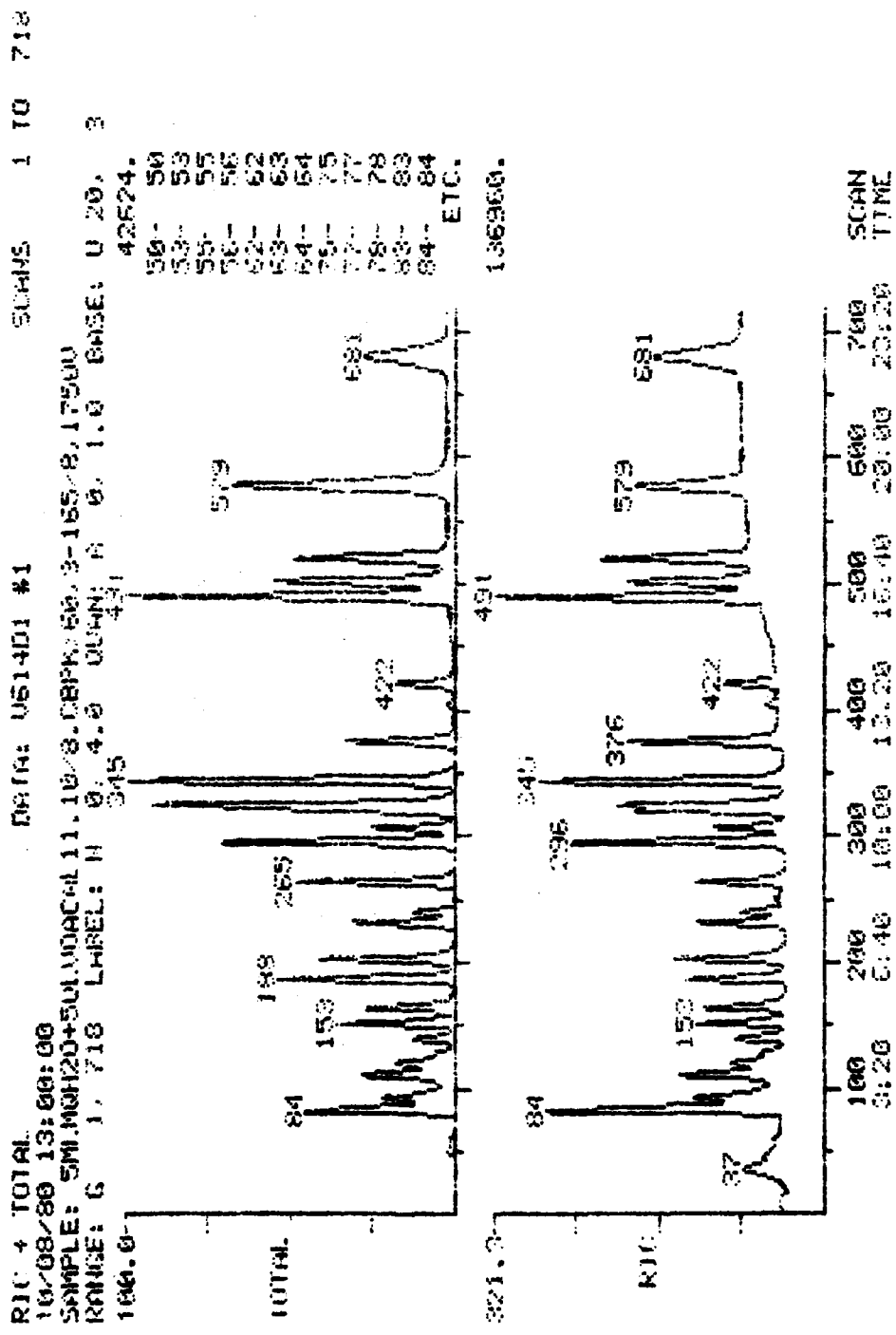
	Method Reference Standard ^a			Raw Wastewater Spike ^b		
	\bar{P}	Sp	% Sp	\bar{P}	Sp	% Sp
Chloromethane	104 ^c	20	19	-	-	-
Dichlorodifluoromethane	-	-	-	-	-	-
Bromomethane	93	31	34	114	1	6
Vinyl chloride	72 ^b	4 ^b	64	122	97	80
Chloroethane	102	36	35	121	13	11
Methylene chloride	105	18	17	116	5	4
Acrolein	114 ^b	17	15	134	15	12
Trichlorodifluoromethane	71	46	65	118	8	/
Acrylonitrile	110	14	13	124	19	16
1,1-Dichloroethylene	90	22	25	108	9	8
1,1-Dichloroethane	111	20	18	117	4	3
Trans-1,2-dichloroethylene	104	13	13	105	6	6
Chloroform	107	20	19	122	16	13
1,2-Dichloroethane	104	11	11	102	9	9
1,1,1-Trichloroethane	105	50	19	110	17	16
Carbon tetrachloride	94	22	24	98	21	21
Bromodichloromethane	102	10	10	96	16	17
1,2-Dichloropropane	102	7	7	90	8	8
Trans-1,3-dichloropropylene	105	14	13	102	2	2
Trichloroethylene	104	13	12	102	21	20
Benzene	104	6	6	110	8	7
Cis-1,3-dichloropropylene	101	19	19	93	4	4
Dibromochloromethane	106	7	/	92	16	18
1,1,2-Trichloroethane	110	9	8	107	15	14
Bromoform	104	17	16	83	20	24
1,1,2,2-Tetrachloroethane	105	11	11	108	10	9
1,1,2,2-Tetrachloroethylene	98	30	30	94	24	26
Toluene	103	15	15	102	16	16
Chlorobenzene	104	16	15	102	16	16
Ethyl benzene	96	22	23	98	20	20

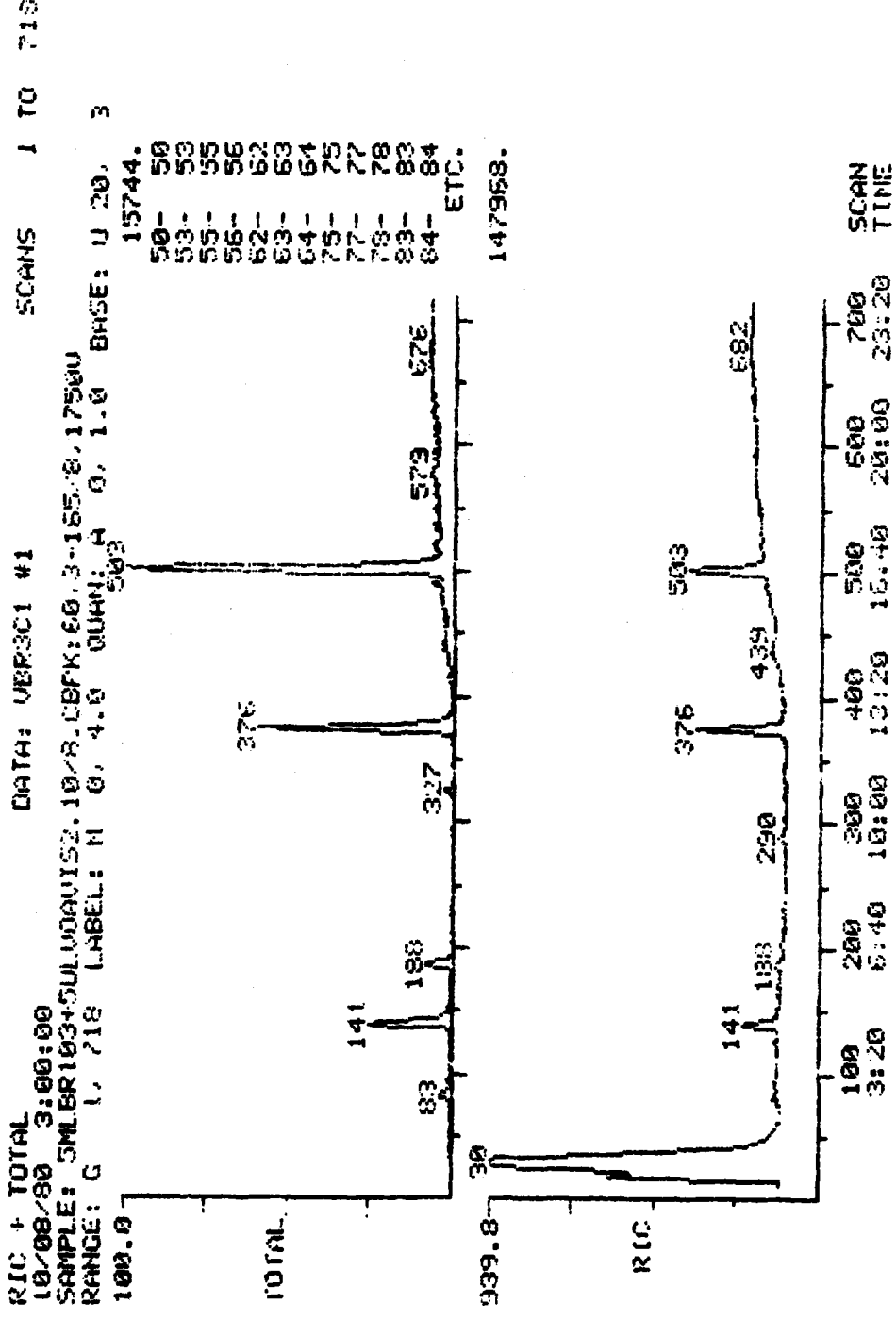
(n) Based on 5 data points, spiked concentrations = 10 $\mu\text{g/L}$ and 25 $\mu\text{g/L}$.

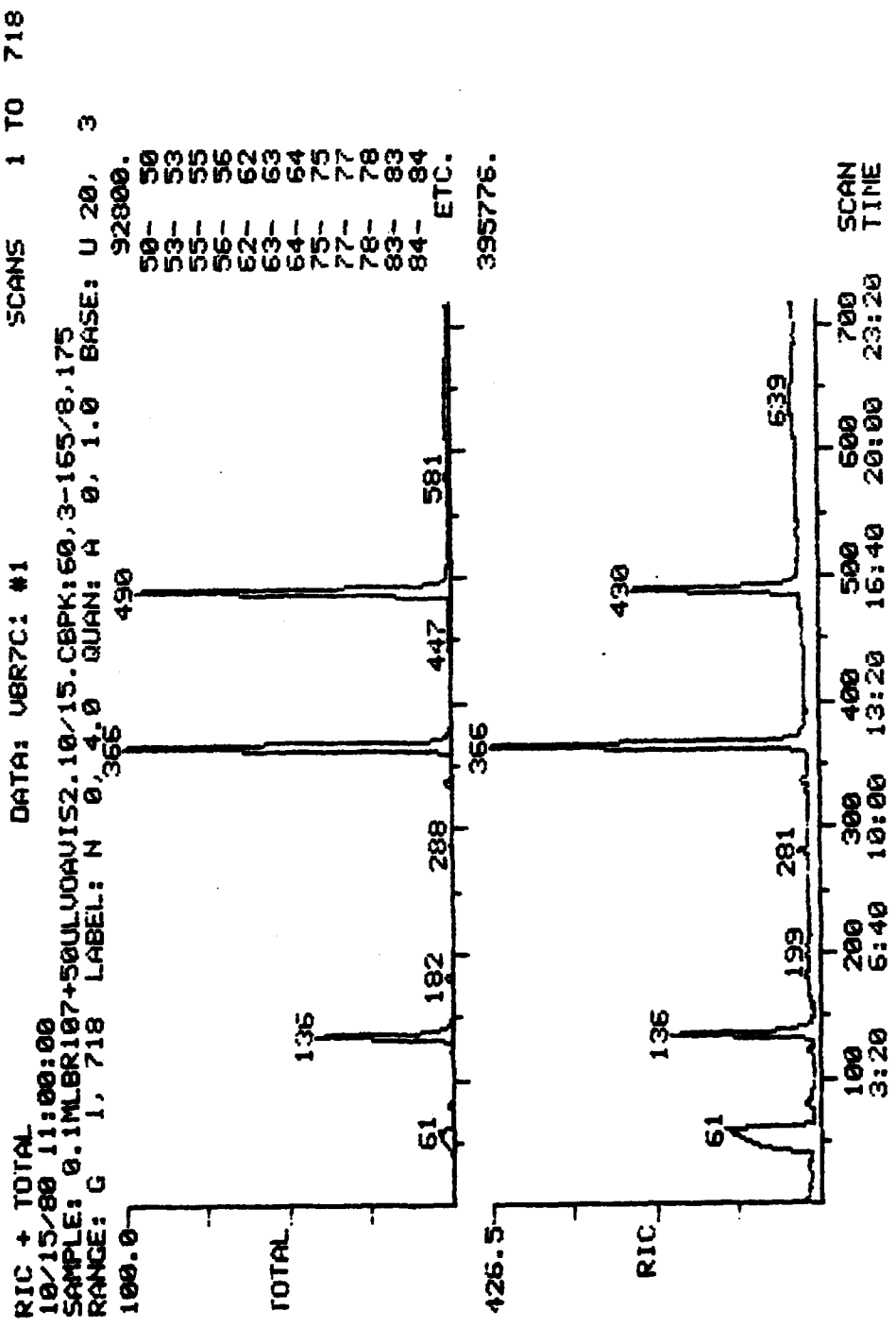
(b) Based on 4 data points, spiked concentrations = 10 $\mu\text{g/L}$ and 25 $\mu\text{g/L}$.

(c) Based on 2 data points, spiked concentrations = 10 $\mu\text{g/L}$ and 25 $\mu\text{g/L}$.

See text page 5, for explanation of Statistics.







APPENDIX B

**Method for the Analyses of Priority Pollutant Acids
in Water**

Priority Pollutant Acids Data

ANALYSIS OF ACID PRIORITY POLLUTANTS
IN WATER

APPLICATION:

METHOD USED TO DETERMINE CONCENTRATIONS OF THE FOLLOWING COMPOUNDS IN WATER SAMPLES

2-CHLOROPHENOL
2-NITROPHENOL
PHENOL
2,4-DIMETHYLPHENOL
2,4-DICHLOROPHENOL
2,4,6-TRICHLOROPHENOL
4-CHLORO-3-CRESOL
2,4-DINITROPHENOL
4,6-DINITRO-2-CRESOL
PENTACHLOROPHENOL
4-NITROPHENOL

- A. TESTED CONCENTRATION RANGE; 5 to 100 UGL FOR EACH
- B. SENSITIVITY; SEE FEDERAL REGISTER 12/3/79 VOL. 4, NO. 233
- C. DETECTION LIMIT;

2-CHLOROPHENOL	33 UGL
2-NITROPHENOL	25 UGL
PHENOL	19 UGL
2,4-DIMETHYLPHENOL	48 UGL
2,4-DICHLOROPHENOL	26 UGL
2,4,6-TRICHLOROPHENOL	22 UGL
4-CHLORO-3-CRESOL	22 UGL
2,4-DINITROPHENOL	73 UGL
4,6-DINITRO-2-CRESOL	45 UGL
PENTACHLOROPHENOL	16 UGL
4-NITROPHENOL	58 UGL
- D. INTERFERENCES; SEE FEDERAL REGISTER 12/3/79 VOL 4, NO. 233
- E. ANALYSIS RATE; AFTER TUNING OF INSTRUMENT, ONE ANALYST CAN ANALYZE 8 SAMPLES IN A 8-HOUR DAY.

CHEMISTRY:

C₆H₅ClO 2-CHLOROPHENOL
CAS RN 95-57-8
MELTING PT 8C BOILING PT 175-176C

C6H5NO₃ 2-NITROPHENOL
CAS RN 88-75-5
MELTING PT 44-45C BOILING PT 214-216C

C6H5OH PHENOL
CAS RN 108-95-2
MELTING PT 39.5-41.5C BOILING PT 181C

C8H10O 2,4-DIMETHYLPHENOL
CAS RN 105-67-9
MELTING PT 25.4-26.0C BOILING PT 212C

C6H4Cl₂O 2,4-DICHLOROPHENOL
CAS RN 120-83-2
MELTING PT 42-43C BOILING PT 209-210C

C6H₃Cl₃O 2,4,6-TRICHLOROPHENOL
CAS RN 88-06-2
MELTING PT 64-66C BOILING PT 246C

C₇H₇ClO 4-CHLORO-3-CRESOL
CAS RN 59-50-7
MELTING PT 65-68C BOILING PT 235C

C₆H₄N₂O₅ 2,4-DINITROPHENOL
CAS RN 51-28-5
MELTING PT 106-108C

C₇H₆N₂O₅ 4,6-DINITRO-2-CRESOL
CAS RN 534-52-1
MELTING PT 87.5C

C₆Cl₅OH PENTACHLOROPHENOL
CAS RN 87-86-5
MELTING PT 188-191C BOILING PT 309-310C

C₆H₅NO₃ 4-NITROPHENOL
CAS RN 100-02-7
MELTING PT 112-114C BOILING PT 279C

USE CAUTION IN HANDLING. POTENTIAL TOXIC INHALATION, AND SKIN ABSORPTION HAZARDS EXIST.

APPARATUS:

- A. INSTRUMENTATION;
GAS CHROMATOGRAPH - FINNIGAN 4000 GAS CHROMATOGRAPH/MASS SPECTROMETER WITH A 6110 DATA SYSTEM

B. PARAMETERS:

COLUMN-1% SP 2250 on 100/120 SUPELCOPORT,

6 FT X 2 MM ID GLASS COLUMN

GAS FLOW - HELIUM GAS AT 30 ML/MIN

TEMPERATURE - HELD AT 50C FOR 4 MINUTES

PROGRAMMED AT 265C AT 8C/MINUTE

HELD CONSTANT AT 265C FOR 30

MINUTES

INJECTOR VOLUME - 2 UL

DETECTOR - MASS SPECTROMETER SET TO ACQUIRE

ELECTRON IMPACT DATA OVER THE MASS

RANGE 41-425 AMV.

SCANNING EVERY 3 SECONDS

RETENTION TIMES - SEE FEDERAL REGISTER

12/3/79 VOL. 4, NO. 233

C. HARDWARE/GLASSWARE; SEE FEDERAL REGISTER

12/3/79 VOL. 4, NO. 233

D. CHEMICALS; SEE FEDERAL REGISTER 12/3/79

VOL. 4, NO. 233

STANDARDS;

A. CALIBRATION STANDARDS; PREPARE STOCK SOLUTION
BY MIXING 12.5 MG OF EACH OF THE FOLLOWING ACIDS
AND DILUTING TO 25 ML IN METHANOL:

2-CHLOROPHENOL

4-CHLORO-3-CRESOL

2-NITROPHENOL

2,4-DINITROPHENOL

PHENOL

4,6-DINITRO-2-CRESOL

2,4-DIMETHYLPHENOL

PENTACHLOROPHENOL

2,4-DICHLOROPHENOL

4-NITROPHENOL

2,4,6-TRICHLOROPHENOL

FINAL CONCENTRATION IS 0.5 MG/ML

PREPARE WORKING CALIBRATION STANDARDS USING
METHYLENE CHLORIDE AS FOLLOWS:

CAL	<u>UL STOCK/2ML CH₂CL₂</u>	<u>FINAL CONC.</u>
A	20 UL	5 UG/ML
B	40 UL	10 UG/ML
C	100 UL	25 UG/ML
D	200 UL	50 UG/ML
E	300 UL	75 UG/ML

10 UL INTERNAL STANDARD SPIKING SOLUTION AND
10 UL D₁₀ ANTHRACENE WERE ADDED TO ALL CALIBRA-
TION STANDARDS AND SAMPLES.

THE INTERNAL STANDARD SPIKING SOLUTION WAS PREPARED AS FOLLOWS;

53.3 MG OCTAFLUOROBIPHENYL AND 51.45 MG 2,4-DIBROMOPHENOL WERE ADDED TO 50 ML VOLUMETRIC FLASK AND DILUTED TO MARK IN METHANOL

FINAL CONCENTRATIONS

1.066 MG/ML OCTAFLUOROBIPHENYL
1.029 MG/ML 2,4-DIBROMOPHENOL

D10 ANTHRACENE STOCK WAS PREPARED AS FOLLOWS:
DILUTE 50 MG D10 ANTHRACENE TO 50 ML IN METHYLENE CHLORIDE. FINAL CONCENTRATION IS 1.0 MG/ML

B. CONTROL SPIKES: A LITER OF NATURAL WATER WAS SPIKED WITH THE ACID STOCK SOLUTION (0.5 MG/ML) AS FOLLOWS:

QC	UL STOCK/L	FINAL CONC.
1	BLANK	
2	10 UL	5 UG/L
3	20 UL	10 UG/L
4	40 UL	20 UG/L
5	100 UL	50 UG/L
6	200 UL	100 UG/L

IN ADDITION, QUALITY CONTROL SAMPLES (1L) WERE SPIKED WITH 10 UL INTERNAL STANDARD SOLUTION. PRIOR TO ANALYSIS ALL 2ML EXTRACTS WERE SPIKED WITH 10 UL D10 ANTHRACENE STOCK

PROCEDURE

SEE FEDERAL REGISTER 12/3/79 VOL. 44, NO. 233

CALCULATIONS:

CALCULATE UGL FOR EACH SAMPLE FROM DAILY CALIBRATION CURVE. USING CALCULATED CONTROL SPIKE CONCENTRATIONS, PLOT UGL ADDED VERSUS UGL FOUND USING DETECTION LIMIT TAPE SUPPLIED BY USATHAMA. CORRECT ALL FIELD SAMPLE CONCENTRATIONS USING THE SLOPE OF THE LINEAR REGRESSION LINE.

REFERENCES:

FEDERAL REGISTER 12/3/79 VOL. 44, NO. 233.
PAGES 69540-69547

TABLE B-1

QC NATURAL WATER SPIKES^a

Task R902.35.08

PRIORITY POLLUTANT CHEMICAL ANALYSIS ($\mu\text{g/L}$)
ACIDS

SPIKE LEVEL	-	+5	+10	+20	+50	+100	R	x(d)
2-Chlorophenol	ND	4.8	7.9	19.9	54.4	87.0	.991	33
2-Nitrophenol	ND	3.4	5.3	15.4	46.4	79.8	.995	25
Phenol	ND	3.0	5.5	13.5	38.4	68.0	.997	19
2,4-Dimethylphenol	ND	1.9	5.1	16.1	42.6	63.4	.982	48
2,4-Dichlorophenol	ND	4.4	7.0	18.3	52.0	87.8	.994	26
2,4,6-Trichlorophenol	ND	4.2	6.1	16.8	49.2	86.0	.996	22
4-Chloro-3-cresol	ND	3.8	5.6	15.6	45.2	78.8	.996	22
2,4-Dinitrophenol	ND	-	0.1	1.1	10.5	54.6	.961	73
4,6-Dinitro-2-cresol	ND	0.2	0.4	2.6	16.9	57.0	.983	45
Pentachlorophenol	ND	3.0	5.0	13.0	35.0	81.0	.998	16
4-Nitrophenol	ND	2.2	1.8	5.5	13.2	48.6	.974	58

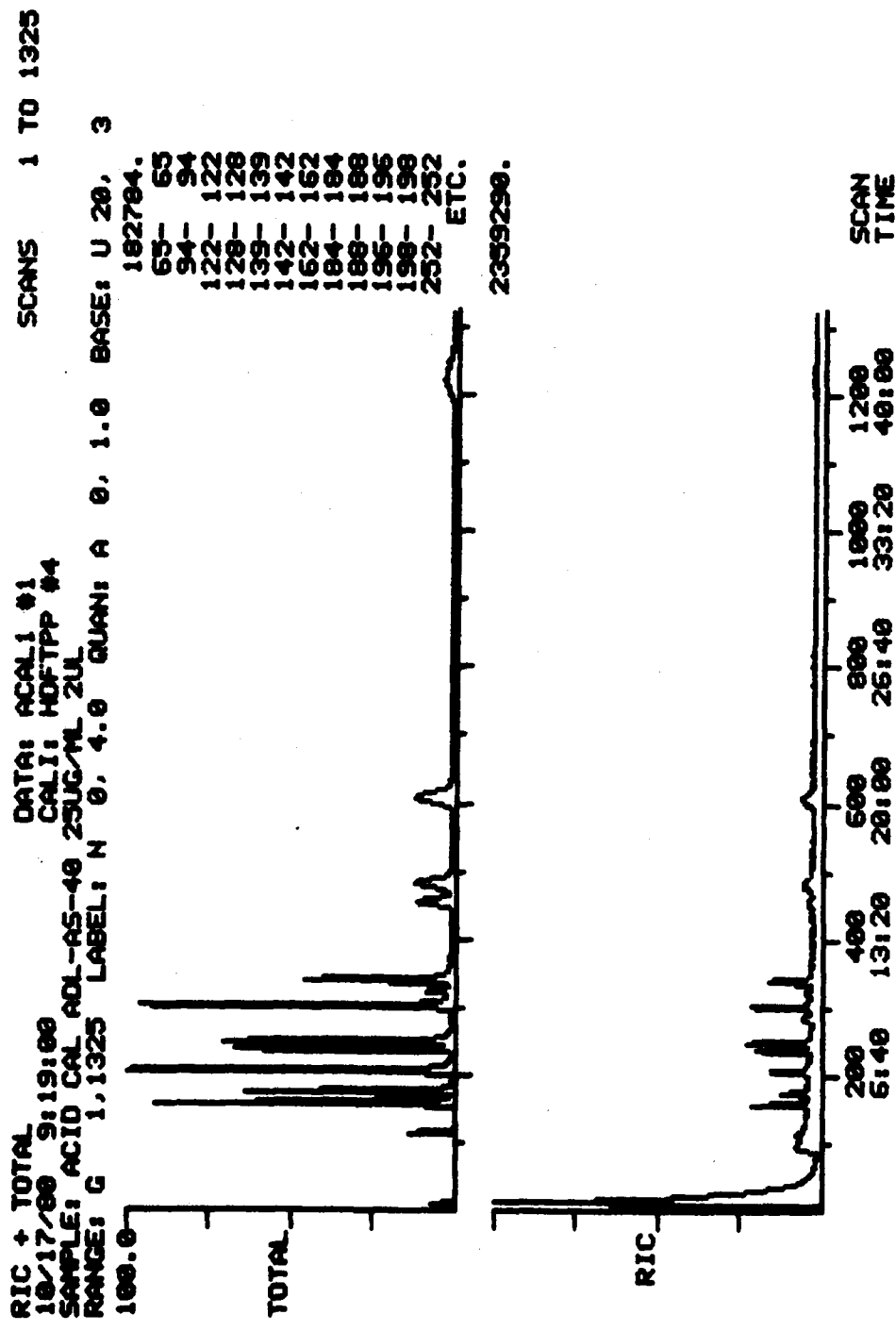
^aField sample from uncontaminated background well bordering a U.S. Army installation site.

ND = none detected

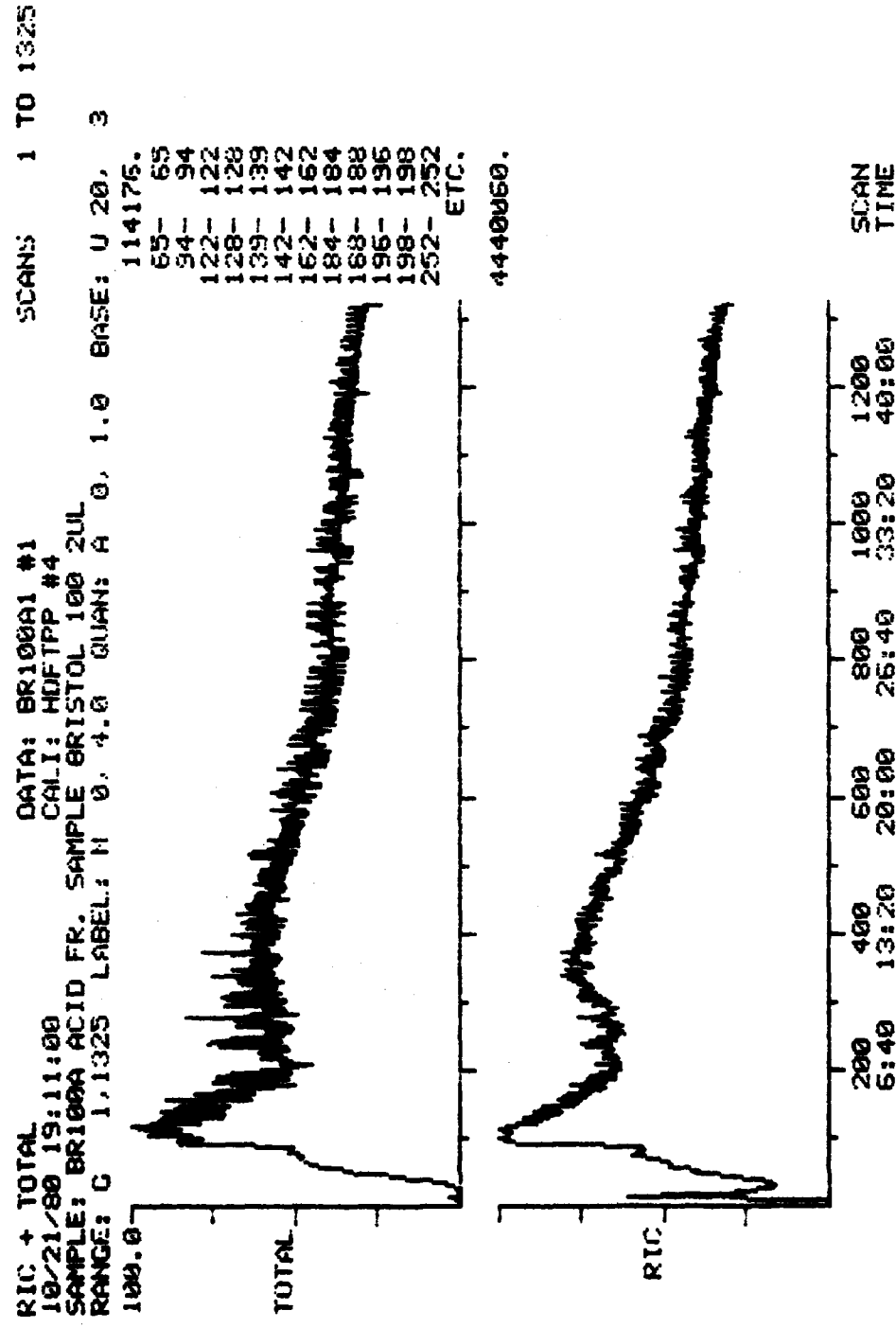
TABLE B-2
EPA QUALITY CONTROL DATA SUMMARY
ACIDS

	Method Reference Standard ^a			Raw Wastewater Spike ^a			
	P	Sp	% Sp	P	Sp	% Sp	
2-Chlorophenol	84	13	16		97 ^b	9	10
2-Nitrophenol	84	7	9		99 ^b	2	2
Phenol	42	11	25		61	24	40
2,4-Dimethylphenol	78	11	14		71 ^b	31	44
2,4-Dichlorophenol	89	12	13		96	18	19
2,4,6-Trichlorophenol	98	12	12		108 ^b	14	13
4-Chloro-3-cresol	86	10	11		82	39	48
2,4-Dinitrophenol	69	9	14		79	2	3
4,6-Dinitro-2-cresol	84	11	13		74	17	22
Pentachlorophenol	79	13	17		64	29	45
4-Nitrophenol	33	6	18		32	3	10

- (a) Based on 4 data points, spiked concentration = 50 µg/L
 The raw waste water spikes were ground water samples associated with a municipal land fill. The data was generated for an EPA sponsored program during the same time period the Bristol samples were analyzed
- (b) Based on 3 data points, spiked concentration = 50 µg/L
 See text, page 5, for explanation of statistics



Acid Calibration Mix



BR 100 Acid Fraction

APPENDIX C

**Method for the Analysis of Priority Pollutant Base/
Neutrals on Water**

Priority Pollutant Base/Neutral Data

ANALYSIS OF BASE/NEUTRALS IN WATER

APPLICATION

METHOD USED TO DETERMINE THE CONCENTRATIONS OF THE FOLLOWING COMPOUNDS:

1,3 DICHLOROBENZENE
1,4 DICHLOROBENZENE
1,2 DICHLOROBENZENE
HEXACHLOROETHANE
BIS(2-CHLOROETHYL)ETHER
NITROSODI-N-PROPYLAMINE
NITROBENZENE
HEXACHLOROBUTADIENE
1,2,4-TRICHLOROBENZENE
BIS(2-CHLOROETHOXY)METHANE
NAPHTHALENE
ISOPHORONE
HEXACHLOROCYCLOPENTADIENE
2-CHLORONAPHTHALENE
ACENAPHTHYLENE
ACENAPHTHENE
2,6-DINITROTOLUENE
4-CHLOROPHENYL PHENYL ETHER
FLUORENE
2,4-DINITROTOLUENE
DIETHYL PHTHALATE

1,2-DIPHENYLHYDRAZINE
N-NITROSODIPHENYLAMINE
HEXACHLOROBENZENE
4-BROMOPHENYL PHENYL ETHER
ANTHRACENE
PHENANTHRENE
DI-N-BUTYL PHTHALATE
FLUORANTHENE
PYRENE
BIS(2-ETHYLHEXYL)PHTHALATE
DI-N-OCTYL PHTHALATE
CHRYSENE
BENZO(A)ANTHRACENE
BENZO(A)PYRENE
INDENO(1,2,3-C,D)PYRENE
DIBENZO(A,H)ANTHRACENE
BENZO(G,H,I)PERYLENE

- A. TESTED CONCENTRATION RANGE: 4-100 µg/L
- B. SENSITIVITY: SEE FEDERAL REGISTER
- C. DETECTION LIMITS: (UGL, BY HUBAUX + VOS)

1,3-DICHLOROBENZENE	19.5
1,4-DICHLOROBENZENE	17.6
1,2-DICHLOROBENZENE	19.4
HEXACHLOROETHANE	17.1
BIS(CHLORO ETHYL)ETHER	24.8

NITROSODI-N-PROPYLAMINE	5.8
NITROBENZENE	15.0
HEXACHLOROBUTADIENE	22.0
1,2,4-TRICHLOROBENZENE	13.0
BIS(2-CHLOROETHOXY)METHANE	9.5
NAPHTHALENE	26.8
ISOPHORONE	10.3
HEXACHLOROCYCLOPENTADIENE	44.8
2-CHLORONAPHTHALENE	12.0
ACENAPHTHYLENE	14.8
ACENAPHTHENE	13.7
2,6-DINITROTOLUENE	17.5
4-CHLOROPHENYL PHENYL ETHER	16.6
FLUORENE	15.7
2,4-DINITROTOLUENE	36.7
DIETHYL PHTHALATE	24.4
1,2-DIPHENYLHYDRAZINE	15.1
N-NITROSODIPHENYLAMINE	13.5
HEXACHLOROBENZENE	18.8
4-BROMOPHENYL PHENYL ETHER	15.1
ANTHRACENE	11.8
PHENANTHRENE	7.2
DI-N-BUTYL PHTHALATE	40.3
FLUORANTHENE	9.8
PYRENE	7.5

BIS(2-ETHYLHEXYL)PHTHALATE	88.0
DI-N-OCTYL PHTHALATE	54.2
CHRYSENE	11.9
BENZO(A)ANTHRACENE	10.8
BENZO(A)PYRENE	19.8
INDENO(1,2,3-C,D)PYRENE	28.3
DIBENZO(A,H)ANTHRACENE	29.2
BENZO(G,H,I)PERYLENE	28.2

D. INTERFERENCES: SEE FEDERAL REGISTER 12/3/79
VOL. 4, NO. 233

E. ANALYSIS RATE: AFTER SAMPLE EXTRACTION AND
CONCENTRATION ONE ANALYST CAN
ANALYZE ONE SAMPLE PER HOUR

CHEMISTRY

C6H4Cl2 1,3-DICHLOROBENZENE
CAS RN 541-73-1
MELTING PT -24C BOILING PT 172-3C

C6H4Cl2 1,4-DICHLOROBENZENE
CAS RN 106-46-7
MELTING PT 54-56C BOILING PT 173C

C6H4Cl2 1,2-DICHLOROBENZENE
CAS RN 95-50-1
MELTING PT -17C BOILING PT 178C

C2Cl6 HEXACHLOROETHANE
CAS RN 67-72-1
MELTING PT 190-195C

C4H8Cl20 BIS(2-CHLOROETHYL)ETHER
CAS RN 111-44-4
MELTING PT -50C BOILING PT 178C

C₆H₁₄N₂O NITROSODI-N-PROPYLAMINE
CAS RN 621-64-7
MELTING PT 52-53C BOILING PT 312C

C₆H₅N₂O NITROBENZENE
CAS RN 98-95-3
MELTING PT 5-6C BOILING PT 210-211C

C₄Cl₆ HEXACHLOROBUTADIENE
CAS RN 87-68-3
MELTING PT -19C BOILING PT 210-220C

C₆H₃Cl₃ 1,2,4-TRICHLOROBENZENE
CAS RN 120-82-1
MELTING PT 16C BOILING PT 214C

C₅H₁₀C₁₂O₂ BIS(2-CHLOROETHOXY)METHANE

C₁₀H₈ NAPHTHALENE
CAS RN 91-20-3
MELTING PT 81-83C BOILING PT 217.7C

C₉H₁₄O ISOPHORONE
CAS RN 78-59-1
MELTING PT -80C BOILING PT 213-214C

C₅Cl₆ HEXACHLOROCYCLOPENTADIENE
CAS RN 77-47-4
MELTING PT -10C BOILING PT 239C

C₁₀H₇C₁ 2-CHLORONAPHTHALENE
CAS RN 91-58-7
MELTING PT 59.5C BOILING PT 256C

C₁₂H₈ ACENAPHTHYLENE
CAS RN 208-96-8
MELTING PT 88-91C BOILING PT 280C

C₁₂H₁₀ ACENAPHTHENE
CAS RN 83-32-9
MELTING PT 93-95C BOILING PT 279C

C₇H₆N₂O₄ 2,6-DINITROTOLUENE
CAS RN 606-20-2
MELTING PT 64-66C

C₁₂H₉ClO 4-CHLOROPHENYL PHENYL ETHER
CAS RN 7005-72-3

C₁₃H₁₀ FLUORENE
CAS RN 86-73-7
MELTING PT 112-115C BOILING PT 298C

C₇H₆N₂O₄ 2,4-DINITROTOLUENE
CAS RN 121-14-2
MELTING PT 69.5 BOILING PT 300C

C₁₂H₁₄O₄ DIETHYL PHTHALATE
CAS RN 84-66-2
MELTING PT -3C BOILING PT 298-299C

C₁₂H₁₂N₂ 1,2-DIPHENYLHYDRAZINE
CAS RN 122-66-7
MELTING PT 123-126C

C₁₂H₁₀N₂O N-NITROSODIPHENYLAMINE
CAS RN 1689-82-3
MELTING PT 144-145C

C₆Cl₆ HEXACHLOROBENZENE
CAS RN 118-74-1
MELTING PT 227-229C BOILING PT 332 C

C₁₂H₉BrO 4-BROMOPHENYL PHENYL ETHER
CAS RN 101-55-3
MELTING PT 18C BOILING PT 305C

C₁₄H₁₀ ANTHRACENE
CAS RN 120-12-7
MELTING PT 214-217C BOILING PT 340C

C₁₄H₁₀ PHENANTHRENE
CAS RN 85-01-8
MELTING PT 99-101C BOILING PT 336C

C₁₆H₂₂O₄ DI-N-BUTYL PHTHALATE
CAS RN 84-74-2
MELTING PT -35C BOILING PT 340C

C₁₆H₁₀ FLUORANTHENE
CAS RN 206-44-0
MELTING PT 107-110C BOILING PT 384C

C₁₆H₁₀ PYRENE
CAS RN 129-00-0
MELTING PT 149-151C BOILING PT 404C

C₂₄H₃₈O₄ BIS(2-ETHYLHEXYL)PHTHALATE
CAS RN 117-81-7
MELTING PT -50C BOILING PT 384C

C₂₄H₃₈O₄ DI-N-OCTYL PHTHALATE
CAS RN 117-81-7
MELTING PT - 50°C BOILING PT 384°C

C₁₈H₁₂ CHRYSENE
CAS RN 218-01-9
MELTING PT 250-253C BOILING PT 448C

C₁₈H₁₂ BENZO(A)ANTHRACENE
CAS RN 56-55-3
MELTING PT 157-159C BOILING PT 437.6C

C₂₀H₁₂ BENZO(A)PYRENE
CAS RN 50-32-8
MELTING PT 179C

C₂₂H₁₂ INDENO (1,2,3,-C,D)PYRENE
CAS RN 193-39-5
MELTING PT 162.5-164C

C₂₂H₁₄ DIBENZO(A,H)ANTHRACENE
CAS RN 53-70-3
MELTING PT 266C BOILING PT 524C

C₂₂H₁₂ BENZO(G,H,I)PERYLENE
CAS RN 191-24-2
MELTING PT 277-279C BOILING PT >7500C

USE CAUTION IN HANDLING, TOXIC INHALATION
AND SKIN ABSORPTION HAZARDS EXIST.

APPARATUS:

A. INSTRUMENTATION:

FINNIGAN 4000 GAS CHROMATOGRAPH/MASS SPECTROMETER
WITH A 6110 DATA AQUISITION SYSTEM.

B. PARAMETERS:

COLUMN: 15 METER FUSED SILICA CAPILLARY
COLUMN COATED WITH SP2100.

TEMPERATURE PROGRAM: 30°C FOR ONE MINUTE:
HEAT RAPIDLY TO 50°C THEN LINEARLY TO
260°C AT 5°/MIN

HYDROGEN CARRIER GAS: 2 ML/MIN

INJECTOR: 260°C GROB-TYPE SPLITLESS BY HOT
NEEDLE TECHNIQUE

TRANSFER LINE: 200°C

SEPARATOR: 200°C

MASS SPECTROMETER SCAN EI DATA OVER MASS RANGE
OF 41-330 AMU EVERY 1.18 SECS.

C. HARDWARE/GLASSWARE: SEE FEDERAL REGISTER

D. CHEMICALS: SEE FEDERAL REGISTER

STANDARDS

A. CALIBRATION STANDARDS:

STOCK A: WEIGH 20 UG OF EACH BASE-NEUTRAL COMPOUND INTO A 100 ML VOLUMETRIC AND DILUTE TO VOLUME IN MEOH.

INTERNAL STANDARD STOCK: WEIGH 5 MG D10 ANTHRA-CENE INTO A 10 ML VOLUMETRIC AND DILUTE TO VOLUME IN MEOH. ADD 10 UL PER 2 ML FOR EVERY SAMPLE AND CALIBRATION STANDARD IMMEDIATELY BEFORE ANALYSIS.

	<u>UL STOCK ADDED TO 2ML CH₂Cl₂</u>	<u>PPB</u>
CAL 1	20 UL	2
CAL 2	50 UL	5
CAL 3	100 UL	10
CAL 4	200 UL	20
CAL 5	400 UL	40

B. CONTROL SPIKES: PREPARE USING STOCK A AND NATURAL WATER

		<u>Conc. of 2 mL CH₂Cl₂ Extr. (UG/L)</u>
QC 1	BLANK	
QC 2	10 UL/L	1
QC 3	25 UL/L	2.5
QC 4	50 UL/L	4.0
QC 5	100 UL/L	10
QC 6	250 UL/L	20

PROCEDURE: SEE FEDERAL REGISTER 12/3/79. Vol. 44, No. 233

CALCULATIONS:

CALCULATE UGL FOR EACH BASE-NEUTRAL AND EACH SAMPLE FROM DAILY CALIBRATION DATA. USING CONTROL SPIKE DATA, PLOT UGL ADDED VERSUS UGL FOUND BY THE METHOD OF HUBAUX AND VOS USING DL TAPE SUPPLIED BY USATHAMA. CORRECT FIELD SAMPLE CONCENTRATIONS USING THE SLOPE OF THE LINEAR REGRESSION LINE.

REFERENCE:

FEDERAL REGISTER 12/3/79. VOL. 44, NO. 233.
PGS. 69540-69547.

TABLE C-1
 QUALITY CONTROL SAMPLES^a
 TASK R902.35.08
 BASE/NEUTRALS ($\mu\text{g/L}$)

SPIKE SAMPLE	+4	+10	+20	+40	+100	m	R	x(d)
1,3 Dichlorobenzene	-	-	6.3	20.8	55.5	.581	.995	19.5
1,4 Dichlorobenzene	-	0.6	8.1	11.2	54.5	.570	.996	17.6
1,2 Dichlorobenzene	-	0.7	8.4	24.6	57.6	.604	.995	19.4
Hexachloroethane	-	1.8	6.3	21.6	56.7	.590	.996	17.1
Bis(2-chloroethyl) ether	-	-	9.5	43.4	104.2	1.10	.992	24.8
Nitrosodi-n-propylamine	4.0	8.3	14.9	33.2	78.8	.788	.999	5.8
Nitrobenzene	2.3	4.8	11.7	33.5	93.4	.954	.997	15.0
Hexachlorobutadiene	-	-	5.3	21.0	53.6	.564	.994	22.0
1,2,4-Trichlorobenzene	-	2.1	9.6	26.3	65.6	.682	.998	13.9
Bis(2-chloroethoxy) methane	0.4	5.7	13.6	34.7	85.7	.889	.999	9.5
Naphthalene	-	-	11.2	37.4	79.9	.846	.991	26.8
Isophorone	1.3	4.3	13.3	33.4	86.6	.889	.999	10.3
Hexachlorocyclopentadiene ^b	13.3	18.0	28.2	54.5	114.9	4.21	.989	44.8
2-Chloronaphthalene	-	4.1	11.9	32.1	78.5	.812	.998	12.0
Acenaphthylene	1.4	5.6	13.2	32.6	85.2	.878	.997	14.8
Acenaphthene	-	5.6	15.7	38.0	86.4	.892	.998	13.7
2,6-Dinitrotoluene	-	-	13.2	26.7	73.0	.758	.996	17.5
4-Chlorophenyl phenyl ether	-	2.7	12.5	35.5	82.9	.865	.996	16.6
Fluorene	-	3.4	12.8	36.0	84.1	.875	.997	15.7
2,4-Dinitrotoluene	-	-	-	27.8	74.2	.788	.983	36.7
Diethyl phthalate	0.1	3.3	2.6	7.4	18.7	.187	.992	24.4
1,2-Diphenylhydrazine	3.2	6.2	13.5	35.1	101.5	1.03	.997	15.1
N-Nitrosodiphenylamine	-	9.1	16.4	39.6	89.4	.914	.998	13.5
Hexachlorobenzene	0.2	5.0	12.3	32.2	68.4	.707	.996	18.8
4-Bromophenyl phenyl ether	0.2	5.1	13.6	36.3	81.8	.846	.997	15.1

^a Spike levels are 0, 12, 30, 60, 120 ppb.

^b Field sample from uncontaminated background well bordering a U.S. Army installation site.

m = slope

QUALITY CONTROL SAMPLES (Cont'd.)

BASE/NEUTRALS ($\mu\text{g/L}$)

SPIKE SAMPLE	+4	+10	+20	+40	+100	m	R	x(d)
Anthracene ^c	-	2.6	6.0	18.3	37.4	.779	.993	11.8
Phenanthrene ^c	-	2.9	6.9	18.6	42.3	.874	.997	7.2
Di-n-butyl phthalate	0.5	6.8	3.3	10.3	29.1	.284	.980	40.3
Fluoranthene ^c	0.1	3.2	6.7	18.2	39.0	.805	.996	9.8
Pyrene ^c	-	3.0	6.6	17.4	38.9	.802	.997	7.5
Bis(2-ethylhexyl) phthalate ^c	17.1	21.2	26.6	46.2	62.2	.545	.927	88.0
Di-n-octyl phthalate ^c	7.7	10.3	13.4	22.2	27.8	.485	.905	54.2
Chrysene ^c	0.1	3.0	6.8	17.2	34.2	.706	.993	11.9
Benzo(a)anthracene ^c	-	4.7	7.1	15.8	32.8	.665	.994	10.8
Benzo(a)pyrene ^c	2.2	4.0	6.6	16.6	27.6	.552	.981	19.8
Indeno (1,2,3-c,d) pyrene ^c	-	5.5	8.0	15.4	24.3	.486	.963	28.3
Dibenzo (a,h) Anthracene ^c	-	6.2	8.2	16.2	25.4	.506	.961	29.2
Benzo (g,h,i) perylene ^c	-	5.0	7.2	15.4	23.8	.481	.963	28.2

^c Spike levels are 0, 2, 5, 10, 20, 50 ppb.

m = slope

TABLE C-2
SUMMARY OF QUALITY CONTROL DATA

**PRIORITY POLLUTANT CHEMICAL ANALYSIS
BASE/NEUTRALS**

SAMPLE NUMBER	Method Ref. Std. ^a			Raw Waste.			Spike ^b		
	—P	Sp	ZSp				—P	Sp	ZSp
1,3 Dichlorobenzene							60 ^c	16	27
1,4 Dichlorobenzene	68	26	38						
1,2 Dichlorobenzene									
Hexachloroethane	62	25	41				66	38	58
Bis(chloromethyl)ether	-						-		
Bis(2-chloroethyl) ether	70	22	32				65	18	28
Bis(2-chloroisopropyl) ether	d						d		
N-Nitrosodimethylamine	12	10	85				17	11	66
Nitrosodi-n-propylamine	40	47	116				77	17	22
Nitrobenzene	70	12	16				82	11	13
Hexachlorobutadiene	63	25	40				55	11	20
1,2,4-Trichlorobenzene	73	27	36				69	9	14
2-Chloroethyl vinyl ether	-						-		
Bis(2-chloroethoxy) methane	74	21	29				52	19	37
Naphthalene	74	8	10				81	5	6
Isophorone	74	19	26				69	20	29
Hexachlorocyclopentadiene	43	31	73				6	5	87
2-Chloronaphthalene	78	18	23				77	11	14
Acenaphthylene	77	14	18				110	35	32
Acenaphthene	82	13	16				81	10	12
Dimethyl phthalate	16	15	93				18	14	77
2,6-Dinitrotoluene	76	12	16				82	14	18
4-Chlorophenyl phenyl ether	88	35	40				79	15	19
Fluorene	87	21	24				80	11	13
2,4-Dinitrotoluene	77	10	14				73	14	19
Diethyl phthalate	40	30	74				59	15	26
1,2-Diphenylhydrazine	71	7	10				73	12	17
N-Nitrosodiphenylamine	76	20	27				85	26	31
Hexachlorobenzene	74	32	43				74	27	37
4-Bromophenyl phenyl ether	84	14	17				90	23	26

a = Based on 4 data points

b = Based on 3 data points

c = Based on 2 data points

d = Standard not available

See text, page 5, for explanation of statistics.

SUMMARY OF QUALITY CONTROL DATA

(con't)

PRIORITY POLLUTANT CHEMICAL ANALYSIS
BASE/NEUTRALS

SAMPLE NUMBER	Method Ref. Std. ^a			Raw Waste. Spike ^b			
	P	Sp	ZSp	P	Sp	ZSp	
Anthracene	84	7	9		85	19	22
Phenanthrene							
Di-n-butyl phthalate	53	37	69		66	20	31
Fluoranthene	74	17	24		84	24	29
Pyrene	74	16	22		108	48	44
Benzidine	-				-		
Butyl benzyl phthalate	34	23	68		29	14	50
Bis(2-ethylhexyl) phthalate	12	8	62		22	16	72
Di-n-octyl phthalate							
Chrysene	50	39	79		59	20	34
Benz(a)anthracene							
3,3'-Dichlorobenzidine	40	52	128		-		
Benzo(b)fluoranthene	d						
Benzo(k)fluoranthene							
Benzo(a)pyrene	42	46	110		45	16	36
Indeno (1,2,3-c,d) pyrene	34	47	139		-		
Dibenzo (a,h) Anthracene	22	44	200		-		
Benzo (g,h,i) perylene	35	51	146		11	18	173

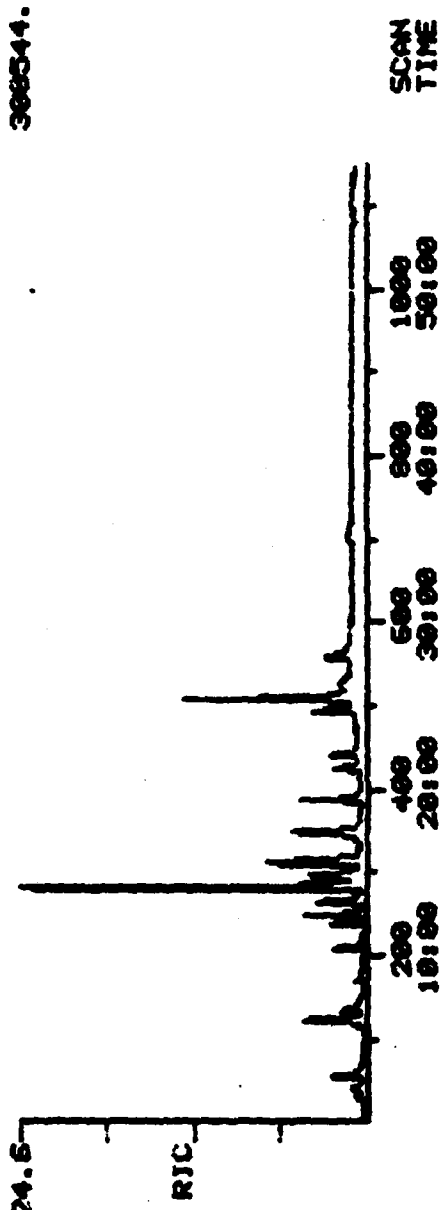
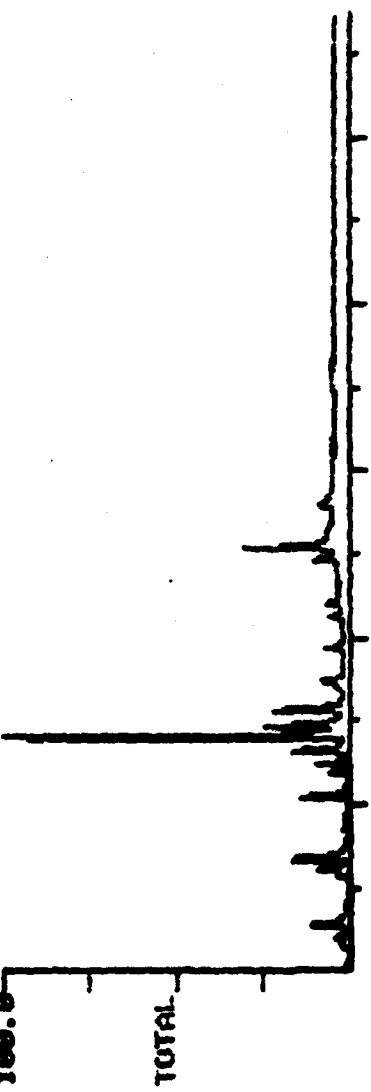
a = Based on 4 data points

b = Based on 3 data points

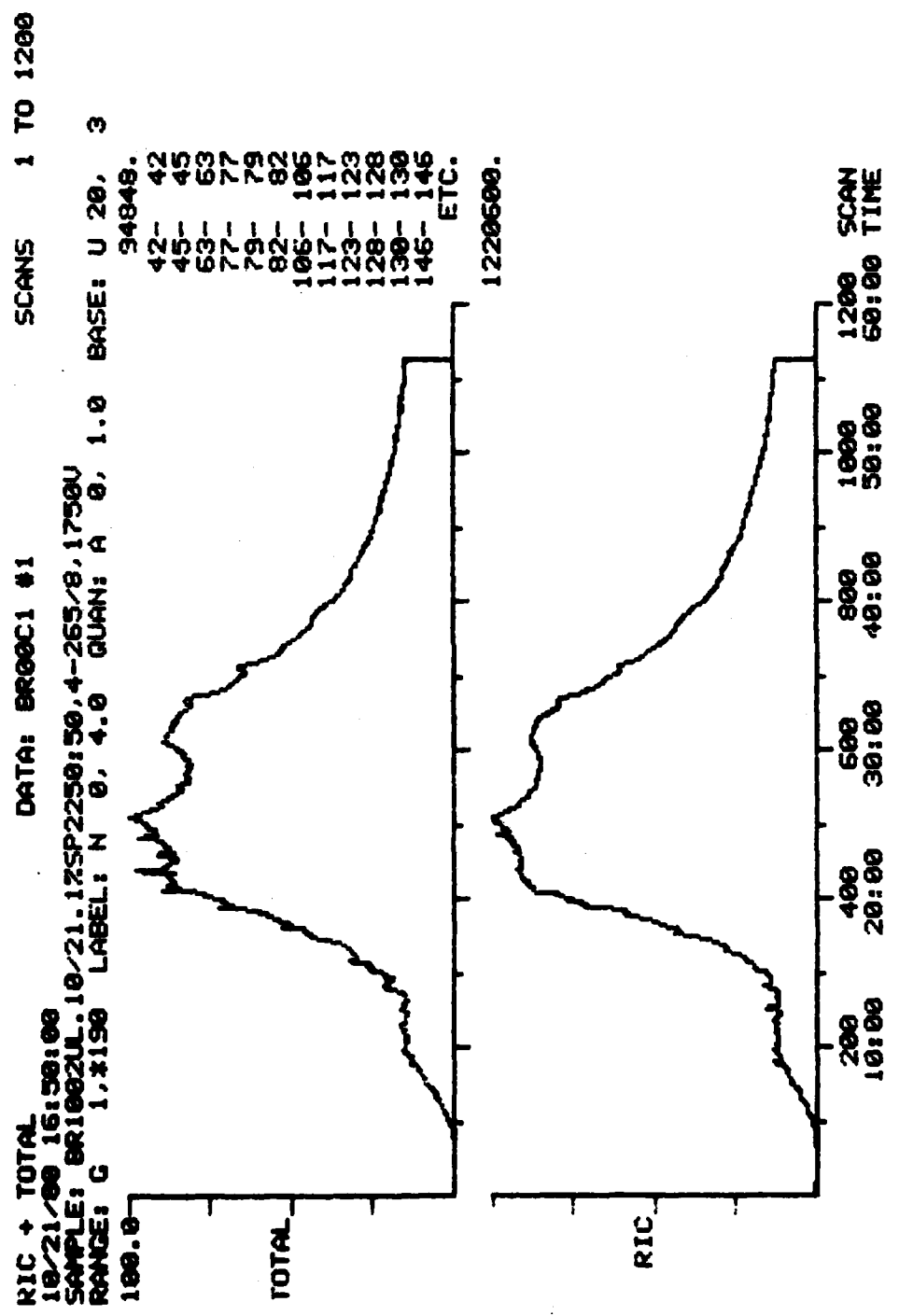
c = Based on 2 data points

d = Standard not available

RIC + TOTAL
 19/21/98 19:38:00
 SAMPLE: 25CAL-214ACN 8549.10/21.17SP2259:59,4:263/8,1736U
 RANGE: G 1,1149 LABEL: N 0, 4.0 BASE: A 0, 1.0
 100.0 78784.
 398544.



Benz - Mental Calibration Mix



APPENDIX D

Method for the Analyses of Pesticides in Water

Priority Pollutant Pesticide/PCB Data

ANALYSIS OF PESTICIDES IN WATER

APPLICATION:

METHOD USED TO DETERMINE THE CONCENTRATIONS OF THE FOLLOWING COMPOUNDS IN WATER SAMPLES:

ALPHA-BHC	ABHC
GAMMA-BHC	GBHC
HEPTACHLOR	HPCL
BETA-BHC	BBHC
DELTA-BHC	DBHC
ALDRIN	ALDRN
HEPTACHLOR EPOXIDE	HPCLE
ENDOSULFAN I	ENDOI
DDE	UDE
DIELDRIN	DLDRN
ENDRIN	ENDRN
DDD	DDD
ENDOSULFAN II	ENDOII
DDT	DDT
PCB 1254	1254
ENDOSULFAN SULFATE	ENDOS

A. TESTED CONCENTRATION RANGE: 1-10 UGL

B. SENSITIVITY:

GC/ECD TUNE REQUIREMENTS: RESPONSE TO 0.05NG
ALDRIN MUST BE AT LEAST 50% FULL SCALE

C. DETECTION LIMIT: (UGL BASED ON HUBAUX + VOS)

ALPHA-BHC	1.3
GAMMA-BHC	0.4
HEPTACHLOR	0.6
BETA-BHC	0.8
DELTA-BHC	0.3
ALDRIN	0.2
HEPTACHLOR EPOXIDE	0.9
ENDOSULFAN I	0.8
DDE	0.4
DIELDRIN	0.4
ENDRIN	0.7
DDD	0.5
ENDOSULFAN II	{ 0.5
DDT	0.7
PCB-1254	0.7
ENDOSULFAN SULFATE	2.5

D. INTERFERENCES: NONE ENCOUNTERED

E. ANALYSIS RATE: AFTER SAMPLE EXTRACTION AND PREPARATION, AND INSTRUMENT CALIBRATION, ONE ANALYST CAN ANALYZE 8 EXTRACTS IN AN 8-HOUR DAY.

CHEMISTRY

ALPHA-BHC C₆ H₆ CL₆
CAS RN 319-84-6
BP 288°C MP 158°C

GAMMA-BHC C₆ H₆ CL₆
CAS RN 58-89-9
BP 323°C MP 113°C

HEPTACHLOR C₁₀ H₅ CL₇
CAS RN 76-44-8
MP 96°C

BETA-BHC C₆ H₆ CL₆
CAS RN 319-85-7
MP 312°C

DELTA-BHC C₆ H₆ CL₆
CAS RN 319-86-8
MP 142°C

ALDRIN C₁₂ H₈ CL₆
CAS RN 309-00-2
MP 104°C

HEPTACHLOR EPOXIDE C₁₁ H₅ OCL₇
CAS RN 1024-57-3

ENDOSULFAN I C₉ H₆ CL₆ O₃S
CAS RN 959-98-8
MP 108°C

DDE C₁₄ H₈ CL₄
CAS RN 72-55-9
MP 88°C

DIELDRIN C₁₂ H₈ CL₆O
CAS RN 60-57-1
MP 176°C

ENDRIN C₁₂ H₈ CL₆O
CAS RN 72-20-8
MP 245°C

DDD C₁₄ H₁₀ CL₄
CAS RN 72-54-8
MP 110°C

ENDOSULFAN II C₉ H₆ CL₆ O₃S
CAS RN 33213-65-9
MP 208°C

DDT C₁₄ H₉ CL₅
CAS RN 50-29-3
MP 107°C

PCB 1254 C₁₂ CL₅ H₅
CAS RN 11097-69-1
MP

ENDOSULFAN SULFATE C₉ H₆ CL₆ O₄S
CAS RN 1031-07-8

USE CAUTION IN HANDLING, POTENTIAL TOXIC
INHALATION AND SKIN ABSORPTION HAZARDS EXIST.

APPARATUS

A. INSTRUMENTATION: HEWLETT PACKARD GAS CHROMATOGRAPH 5840A EQUIPPED WITH AN ELECTRON CAPTURE DETECTOR.

B. PARAMETERS:

COLUMN: 1.5% SP 2250, 1.95% SP 2401, 4MM ID,
6" SPAN, 180 CM.

CARRIER GAS: 5% METHANE, 95% ARGON
TEMPERATURE: 205°C (WHILE RUNNING), 225°C
(BAKING)

RUN TIME: 30 MINUTES

INJECTION PORT TEMPERATURE: 250°C

ECD TEMPERATURE: 300°C

CHART SPEED: 0.50

ZERO: 10.0

ATTENUATION: 2⁺⁸

SLOPE SENSITIVITY: 0.15

AREA REJECTION: 1,000,000

FLOW: 75 ML

AREA REJECTION AT 1.00 MINUTE: 5000

ATTENUATION AT 18 MINUTES: 2⁺⁷

OPTION 3: 1,2

INJECTION VOLUME: 2 UL

C. HARDWARE/GLASSWARE:

KUDERNA-DANISH GLASSWARE INCLUDING THREE
BALL MACRO SNYDER COLUMNS: 500 ML EVAPO-
RATOR FLASK: 10 ML GRADUATED RECEIVER
AMPULE: AMPULE STOPPER.

SEPARATORY FUNNELS - 125, 1000 AND 2000 ML,
WITH TEFLON STOPCOCKS.

CHROMAFLEX COLUMN, 400 MM LONG X 19 MM
(KONTES K-420540-9011)

CENTRIFUGE BOTTLES - 250 ML

CENTRIFUGE

GRADUATED CYLINDERS - 100, 250, 500,
1000 ML

BEAKERS - 100, 400, 600 ML

SYRINGES - 10 - 500 UL

AMBER GLASS BOTTLES WITH TEFLON LINED CAPS,
500 ML AND 1-L

GLASS WOOL - METHYLENE CHLORIDE EXTRACTED

FLORISIL - PR GRADE 60-100 MESH

D. CHEMICALS:

HEXANE, FISHER BRAND, PESTICIDE GRADE
METHYLENE CHLORIDE, FISHER BRAND, PESTICIDE
GRADE

METHANOL, FISHER BRAND, PESTICIDE GRADE

ETHYL ETHER, FISHER BRAND, PESTICIDE GRADE

PETROLEUM ETHER, FISHER BRAND, PESTICIDE GRADE

SODIUM SULFATE, GRANULAR, ANHYDROUS,
CONDITIONED FOR TWO HOURS AT 550°C

PESTICIDE STANDARDS OBTAINED FROM EPA HEALTH
EFFECTS RESEARCH LABORATORY, ENVIRONMENTAL
TOXICOLOGY DIVISION, RESEARCH TRIANGLE PARK,
NORTH CAROLINA.

TETRACHLOROTETRAHYDRONAPHTHALENE (TCTHN, INTERNAL
STANDARD), EASTMAN CHEM. CO.

A. CALIBRATION STANDARDS:

STOCK A: WEIGH 25 MG OF EACH PESTICIDE INTO
A SINGLE 25 ML VOLUMETRIC AND DILUTE
TO VOLUME WITH MEOH TO MAKE A 1 MG/ML SOLUTION

DILUTION A: TAKE 1 ML STOCK A AND DILUTE TO
10 ML WITH MEOH TO MAKE A 0.1 MG/ML
SOLUTION

INTERNAL STANDARD STOCK: WEIGH 20 MG TCTHN
INTO 100 ML VOLUMETRIC AND DILUTE TO
VOLUME WITH MEOH TO MAKE A 0.2
MG/ML SOLUTION. ADD 10 UL OF INTERNAL
STANDARD TO EVERY 10 ML CALIBRATION
SOLUTION AND SAMPLE IMMEDIATELY
BEFORE ANALYSIS (10 UL/10 ML)

CALIBRATION STANDARDS:

	<u>UG/ML</u>
CAL 1 1 UL DIL. A/10ML HEXANE	0.01
CAL 2 2.5 UL DIL.A/10ML HEXANE	0.025
CAL 3 5.0 UL DIL.A/10ML HEXANE	0.05
CAL 4 7.5 UL DIL.A/10ML HEXANE	0.075
CAL 5 10 UL DIL.A/10ML HEXANE	0.1

B. CONTROL SPIKES: PREPARE USING DILUTION A STOCK, LISTED ABOVE, AND NATURAL WATER:

	<u>UG/ML</u>	<u>PPB*</u>
QC 1-BLANK	-	-
QC 2-10 UL DIL.A/L	0.01	1
QC 3-25 UL DIL.A/L	0.025	2.5
QC 4-50 UL DIL.A/L	0.05	5
QC 5-75 UL DIL.A/L	0.075	7.5
QC 6-100 UL DIL.A/L	0.1	10

*BASED on 100 ML FINAL VOLUME

PROCEDURE:

SEE REF. 1 FOR LIQUID-LIQUID EXTRACTION, FOR KUDERIA-DANISH CONCENTRATION, FOR FLORISIL PREPARATION AND SAMPLE CLEANUP, AND FOR SULFUR REMOVAL PROCEDURES.

CALCULATIONS:

CALCULATE UGL FOR EACH PESTICIDE AND EACH SAMPLE FROM DAILY CALIBRATION DATA. USING CONTROL SPIKE DATA, PLOT UGL ADDED VERSUS UGL FOUND BY THE METHOD OF HUBAUX AND VOS, USING DL TAPE SUPPLIED BY USATHAMA. CORRECT FIELD SAMPLE CONCENTRATIONS USING THIS LINE AND LINEAR REGRESSION ANALYSIS.

REFERENCE

1. U.S. EPA EVALUATION OF PROTOCOLS FOR PESTICIDES AND PCB'S IN RAW WASTEWATER. EPA-600/2-79-166. PP. 87-99.

TABLE D-1
Quality Control Samples
Task R902.35.08
Pesticides
 $(\mu\text{g/L})$

Spike Level	+1.0	+2.5	+5.0	+7.5	+10	m	R	x(d)
alpha-BHC	1.39	2.51	4.70	6.77	8.69	.852	.998	1.3
gamma-BHC	0.89	2.14	4.43	6.63	8.67	.873	1.00	0.4
Heptachlor	0.79	2.04	4.24	6.25	8.12	.821	1.00	0.6
beta-BHC	0.77	2.18	4.60	6.74	8.67	.883	.999	0.8
delta-BHC	1.00	2.19	4.43	6.55	8.69	.865	1.00	0.3
Aldrin	0.86	2.07	4.10	6.12	8.25	.820	1.00	0.2
Heptachlor epoxide	0.66	2.13	4.41	6.52	8.38	.856	.999	0.9
Endosulfan I.	0.67	2.09	4.39	6.43	8.33	.850	.999	0.8
DDE	0.83	2.12	4.59	6.92	9.14	.925	1.00	0.4
Dieldrin	.96	2.38	4.97	7.28	9.68	.971	1.00	0.4
Endrin	0.65	2.07	4.25	6.30	8.22	.836	1.00	0.7
DDD	0.68	2.03	4.27	6.38	8.42	.854	1.00	0.5
Endosulfan II								
DDT	0.59	2.27	4.67	7.10	9.49	.966	1.00	0.7
PCB-1254	0.86	2.01	4.40	6.56	9.06	.903	1.00	0.7
Endosulfan sulfate	-	2.55	6.85	9.31	12.2	1.30	.993	2.5

m = slope

R = correlation coefficient

x(d) = detection limit calculated by the method of Hubaux
and Vos

TABLE D-2
EPA QUALITY CONTROL DATA SUMMARY

Pesticides

Method Ref. Std.^a

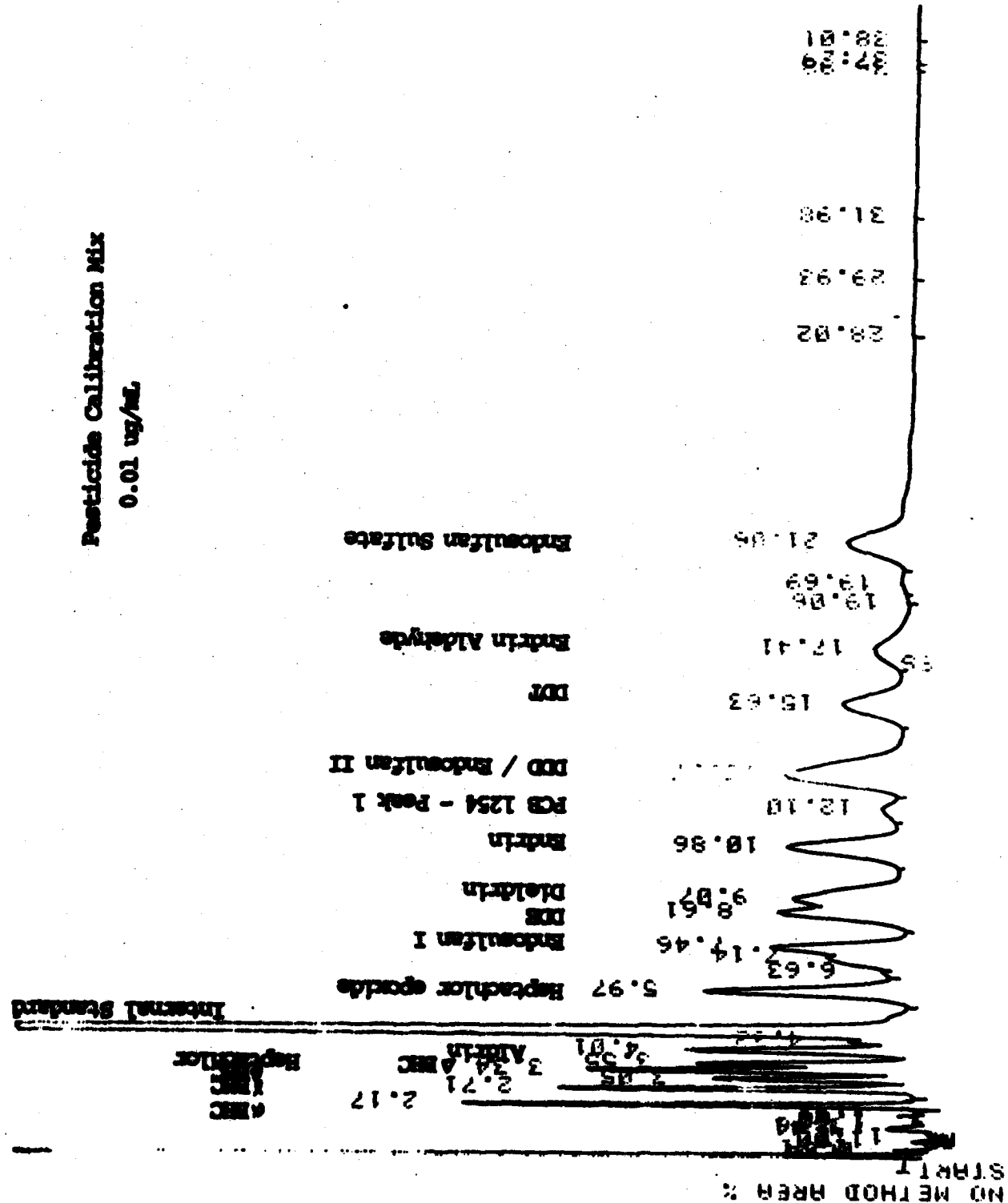
Raw Waste. Spike^b

SAMPLE NUMBER	1	Sp	ZSp				1	Sp	ZSp
alpha-BHC	109	35	32			92	4	4	
gamma-BHC	94	2	2			91	4	4	
Heptachlor	91	1	1			85	4	4	
beta-BHC	94	3	3			88	6	7	
delta-BHC	93	4	4			91	4	5	
Aldrin	91	4	4			83	4	4	
Heptachlor epoxide	92	2	2			85	7	8	
Endosulfan I.	96	3	4			95	14	14	
DDE	100	8	8			91	17	19	
Disdrin	89	8	9			87	20	24	
Endrin	88	2	2			93	8	8	
DDD									
Endosulfan II	90	6	6			88	16	18	
DDT	95	4	4			85	10	11	
PCB-1254	84	7	8			74	7	9	

a = Based on 4 data points

b = Based on 3 data points

See text, page 5, for explanation of statistics.



Astroblar 1016

44.14
43.45

38.92

34.39

31.67

29.58

29.98

29.29

15.51

11.39

19.61

15.38

14.54

13.87

12.19

4.88

4.74

1.14

1.12

1.11

1.11

Archolar 1254

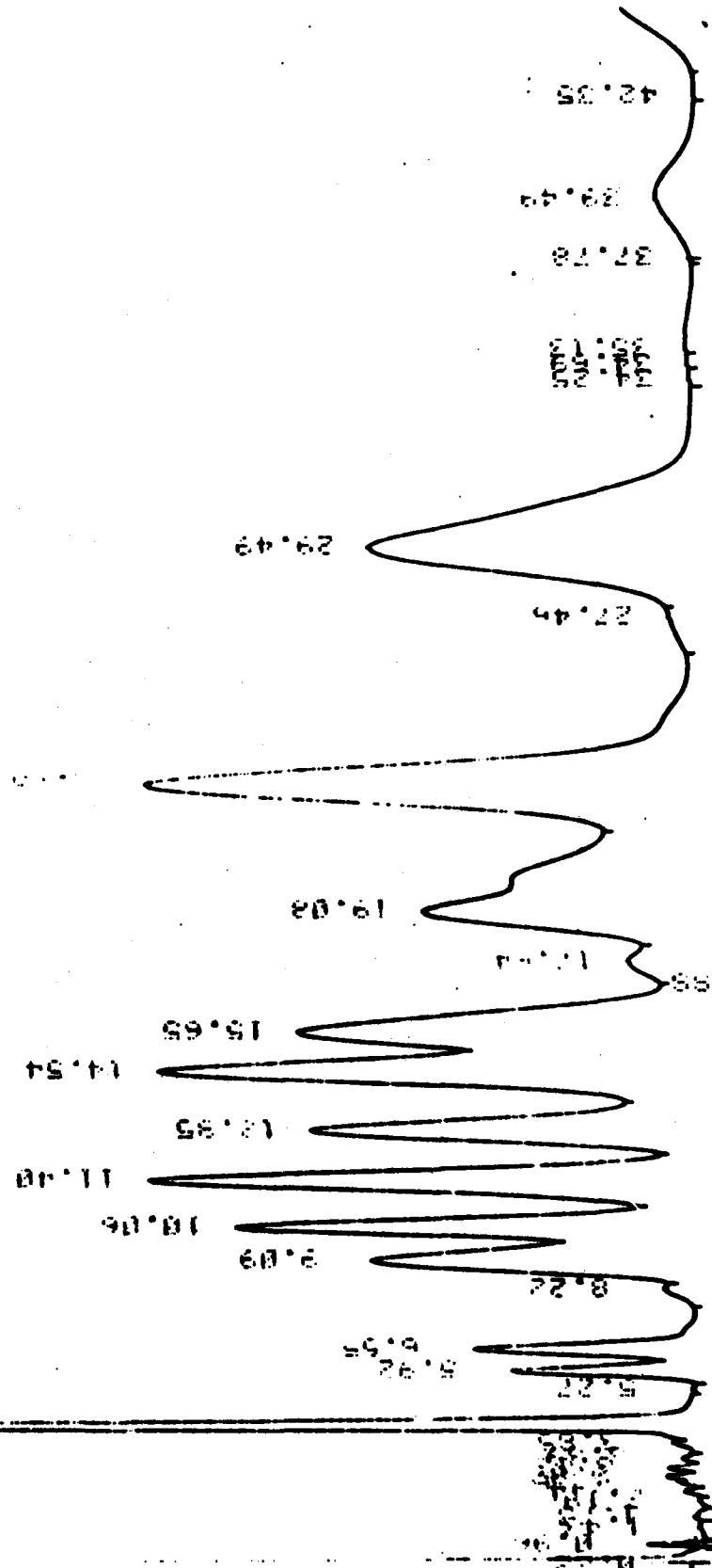
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30.51
29.48
28.45
27.42
26.39
25.36
24.33
23.30
22.27
21.24
20.21
19.18
18.15
17.12
16.09
15.06
14.03
13.00
12.97
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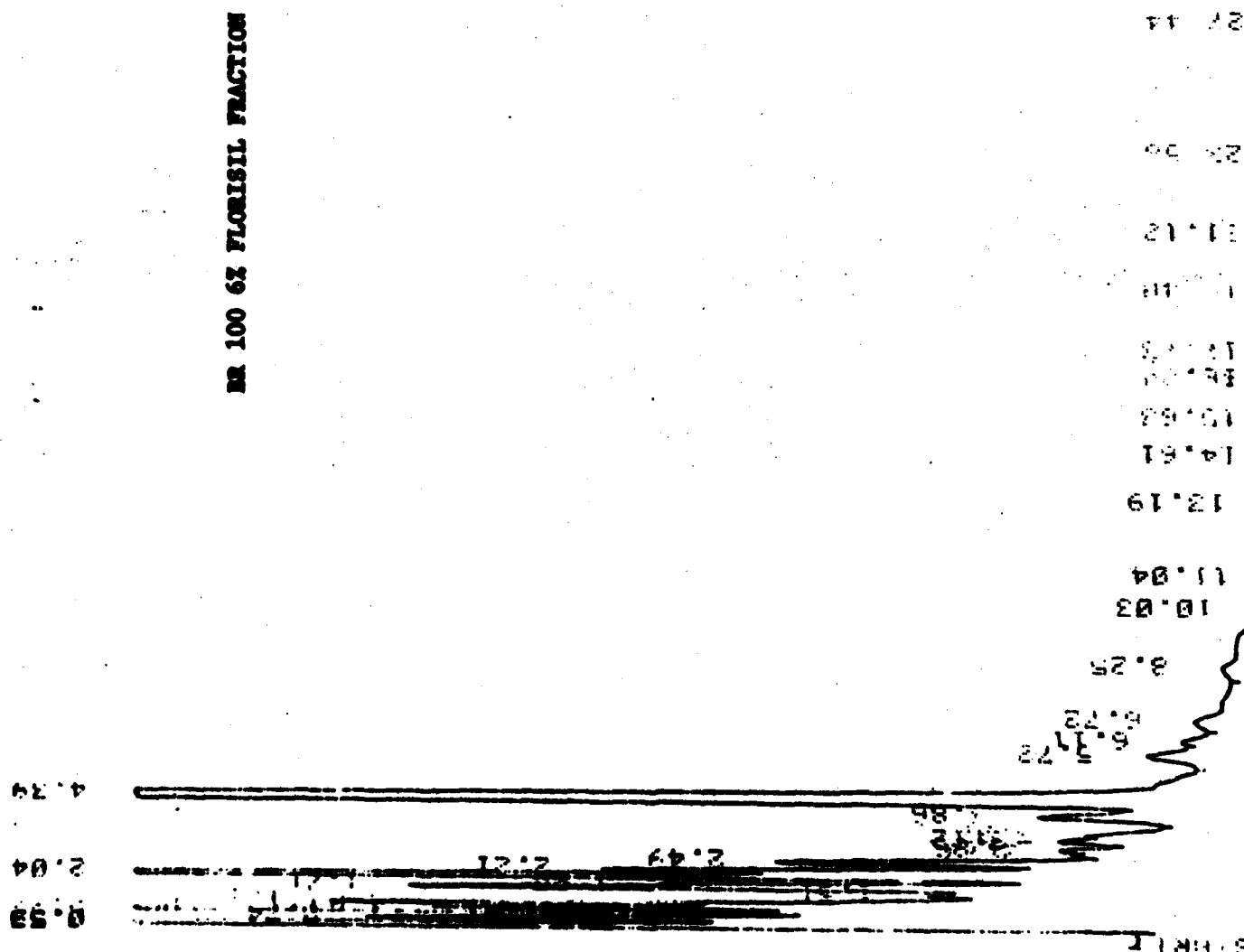
STABIL

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D-12

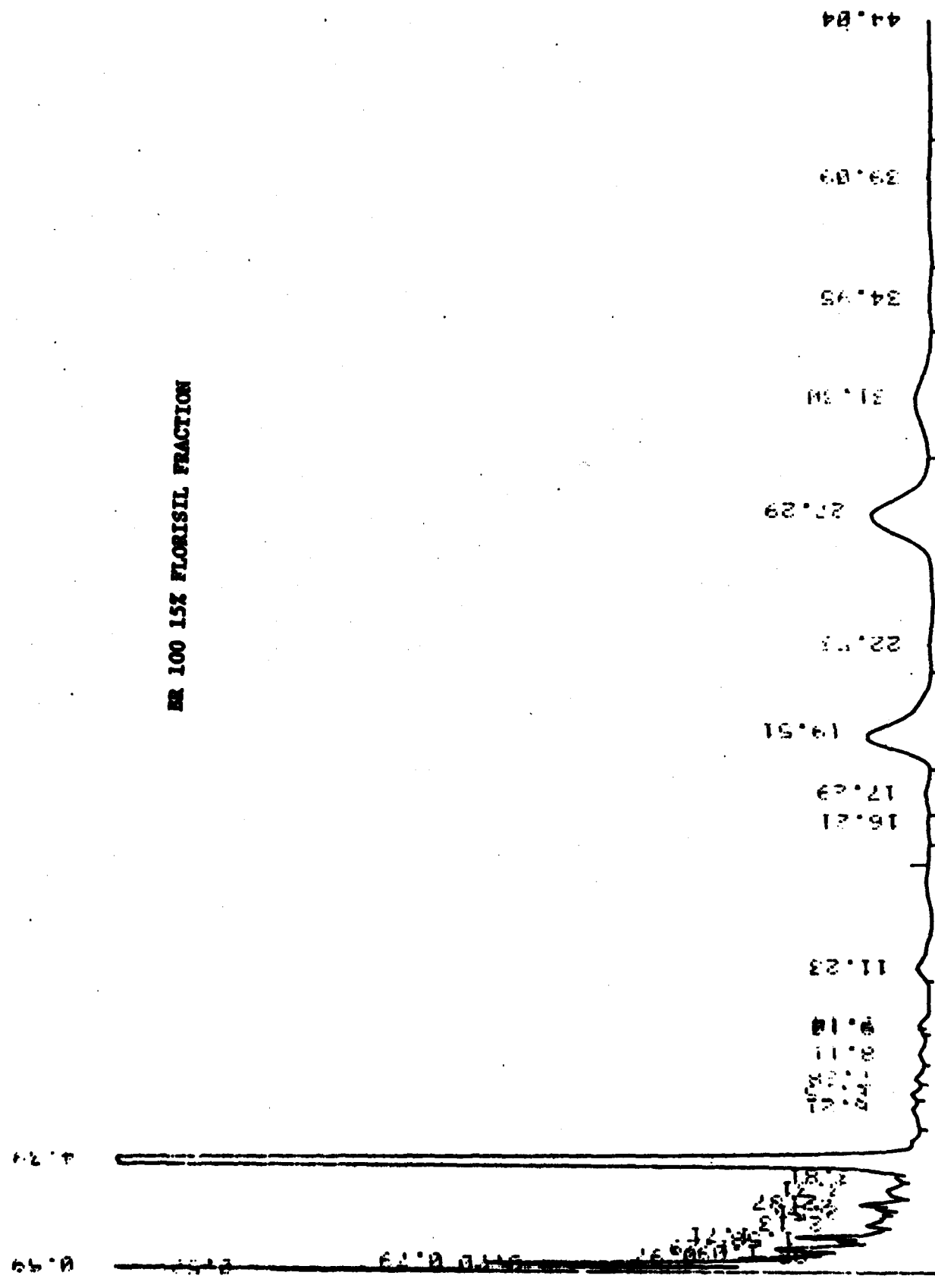
Astrochlor 1260





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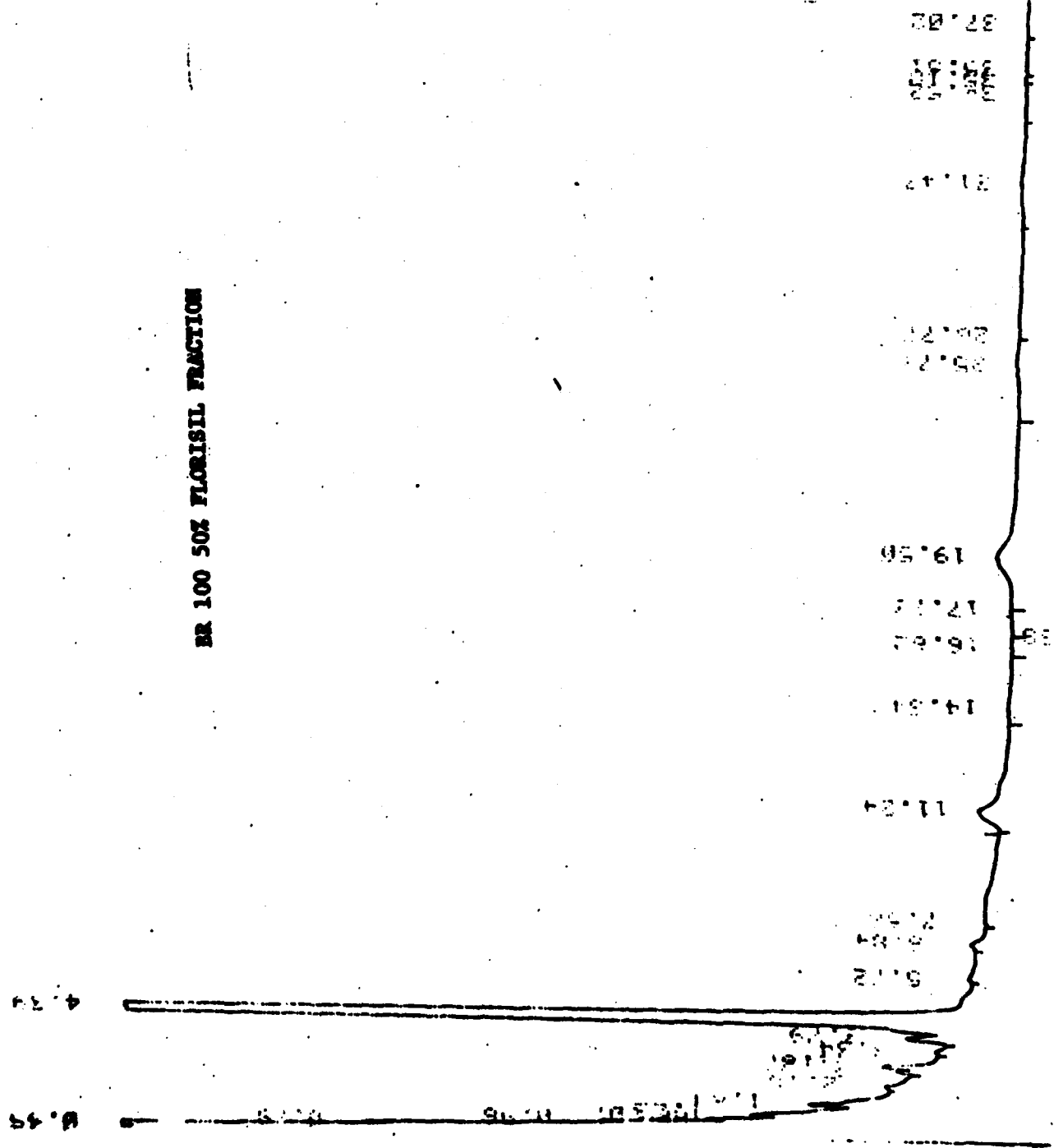
■ 100% FRACTION



D-15

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IN 100 SOZ MORSEIN FRACTION



D-16

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